

Version 1, October, 12th, 2021

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This vacancy booklet is meant for Mexican students intending to enrol in a PhD program abroad, using a <u>LPDP</u>, a <u>KemenDikBud</u> or other PhD scholarship. This booklet gives an overview of PhD vacancies available at Erasmus MC for (candidate) PhD scholarship holders.

For students in biomedical sciences, biomedical engineering, computer science, health sciences, medicine, pharmacy and vet medicine.

Pendahuluan Erasmus MC



Preclinical, clinical & Health Sciences 2016-2020 InCites Clarivate dbase as of Oct. 5th, 202 University or Med School only* publ world impact 24.271 2 55 Erasmus University Rotterdam 25,746 2.52 UCLA DG Med School* 15,863 2.47 Harvard University 139,589 2.37 Stanford University 40,396 2.32 Johns Hopkins University 63,010 2.27 ohns Hopkins Medicine* 22,879 2.27 Harvard Univ Med School* 70,795 2.27 UC San Francisco 47,712 2.22 Yale University 34,241 2.21 UC Los Angeles (UCLA) 37,742 2.21 University of Chicago 16,265 2.13



Erasmus University Medical Center, yang dikenal sebagai Erasmus MC, Sekolah Kedokteran Erasmus University, dan 3 rumah sakit universitasnya terintegrasi di dalam satu kampus dan dipimpin oleh satu direksi. Pusat pendidikan dengan 400 tempat belajar dan 40 ruang kelas untuk sampai 6.000 mahasiswa ini dibuka pada tahun 2012 dan diberi penghargaan untuk arsitekturnya pada tahun 2013. Pada tahun 2018, rumah sakit-rumah sakit tua diganti dengan rumah sakit canggih yang memiliki 1.000 kamar untuk pasien tunggal. Erasmus MC berkomitmen pada penduduk yang sehat dan keunggulan dalam layanan kesehatan melalui riset dan pendidikan (www.erasmusmc.nl).

Perawatan pasien: Erasmus MC, yang hanya puas dengan perawatan terbaik, hanya memiliki kamar pasien tunggal (rumah sakit VIP) untuk mempercepat inovasi kedokteran dan kemampuannya untuk merawat pasien dengan bahan dan prosedur terbaru dan paling inovatif. <u>https://www.youtube.com/watch?v=agYQOLrhmrQ</u>

Riset & Inovasi: Erasmus MC secara konsisten menempati peringkat 13-36 teratas di dunia untuk berbagai bidang klinis dan peringkat 30 teratas untuk ilmu biomedis (<u>US News Clinical Medicine 2021</u>, <u>Nature Index Biomedical Sciences 2019</u>). Hal yang penting adalah dampak makalah risetnya di seluruh dunia dalam ilmu praklinis, klinis, & kesehatan memiliki skor 2,55, yang menempati peringkat atas dunia, persis di atas Harvard (2,37 lihat tabel atas). Tujuan umum riset Erasmus MC adalah menerjemahkan penemuan di laboratorium ke penerapan praktis pada pasien dan mencakup semua bidang mulai dari riset praklinis melalui riset klinis sampai riset ilmu kesehatan.

Pendidikan & Pelatihan: Erasmus MC menawarkan program BSc, MSc, PhD, dan Residensi untuk melatih generasi praktisi dan peneliti medis berikutnya. Erasmus MC adalah salah satu sekolah kedokteran terbesar di Eropa, dengan ~2.500 mahasiswa kedokteran dan antara 220-250 kelulusan PhD/tahun. Dengan 33% mahasiswa kedokterannya yang telah menerbitkan makalah, 70% di luar negeri dan 20% memilih MD-PhD (untuk menjadi dokter klinis dan ilmuwan), <u>pendidikan kedokteran</u> Erasmus MC cukup istimewa. Erasmus MC juga mengharapkan <u>mahasiswa PhD-nya</u> memiliki 4 atau lebih publikasi penelitian (dalam 25% jurnal terakhir dalam bidang penelitiannya) sebelum mengikuti ujian kelulusan. Semua mahasiswa PhD memiliki gelar MSc, MD, atau DVM pada saat mendaftar masuk dan sebagian besar memiliki beasiswa perseorangan atau dibiayai oleh dana hibah penelitian. <u>Program pendidikan inovatif</u>: Erasmus MC dan Delft University of Technology</u> adalah dua universitas pertama di dunia yang menawarkan program BSc-MSc dalam nanobiologi, menjembatani celah antara ilmu hayati & teknologi. Kolaborasi intensif dengan universitas teknologi ini menghasilkan rentang kolaborasi penelitian yang lebih luas dan fokus lebih mendalam pada penerapan langsung di masyarakat. <u>Tingkat pembimbingan</u>: dengan ~750 spesialis kedokteran terdaftar vs ~1.000 residen dan ~1.500 staf sains (ditambah 600 mahasiswa pascadoktoral) vs ~1.250 mahasiswa PhD, kami memiliki salah satu rasio pembimbing terbaik di dunia (mahasiswa PhD memiliki setidaknya dua pembimbing).

Erasmus MC & Eropa: Erasmus MC termasuk ke dalam 10 sekolah kedokteran terbesar di Uni Eropa sebagaimana diukur berdasarkan jumlah publikasi dan jumlah publikasi yang berasal dari penelitian yang didanai EC (yaitu program FP6, FP7 dan Horizon2020) dan Erasmus MC adalah salah satu sekolah kedokteran tersukses di Eropa kontinental dalam Horizon2020 bertema Kesehatan, Perubahan Demografis, & Kesejahteraan (lihat tabel kanan pada hal. 3). Dengan demikian, Erasmus MC adalah pintu gerbang yang menarik untuk menuju ke jejaring penelitian Eropa, yang merupakan manfaat yang diperoleh setelah kelulusan Anda terlepas dari apakah karier Anda di dalam atau di luar Eropa.

Kolaborasi di seluruh dunia

Erasmus MC dikenal dengan kolaborasi jangka panjang dan loyalitas mitranya. Filosofi ini diterjemahkan ke dalam kolaborasi penelitian berkualitas tinggi, sebagaimana ditunjukkan oleh jumlah sitasi rata-rata. Kualitas penelitiannya sering jauh lebih baik daripada kualitas penelitian universitas asing dengan mitra mereka yang lebih terkenal (lihat tabel, di bagian atas halaman) dan hal yang penting dalam kolaborasi penelitian adalah karya yang Anda publikasikan bersama.

Program PhD/S3 di Erasmus MC - gambaran

Memilih universitas untuk program PhD adalah langkah terpenting dalam karier yang berorientasi pada penelitian. Program PhD adalah program pendidikan tertinggi yang ditawarkan oleh universitas dan keluaran (yaitu hasil) pelatihan PhD Anda menentukan langkah berikutnya dalam karier Anda. Karena PhD pada dasarnya adalah program pendidikan & pelatihan penelitian, kualitas publikasi penelitian universitas yang Anda bidik menjadi sangat penting. Kami juga memperhatikan bahwa delegasi universitas Eropa dan non-Eropa selalu menekankan pada pentingnya akses ke dana hibah penelitian Eropa. Jadi, jika Anda ingin mengejar karier dalam konteks internasional, ketahui bahwa Erasmus MC memiliki rekam jejak yang baik dalam kualitas makalah penelitiannya serta dalam memperoleh dana hibah penelitian Eropa (yang disebut dana hibah Horizon2020, bertema Kesehatan, Perubahan Demografis, & Kesejahteraan).

Preclinical, clinical & Health Sciences 2016-2020		
InCites Clarivate dbase as of Oct, 5th, 2021		
University or Med School only*	publ	world impact
Erasmus MC*	24,271	2.55
Erasmus University Rotterdam	25,746	2.52
UCLA DG Med School*	15,863	2.47
Harvard University	139,589	2.37
Stanford University	40,396	2.32
Johns Hopkins University	63,010	2.27
Johns Hopkins Medicine*	22,879	2.27
Harvard Univ Med School*	70,795	2.27
UC San Francisco	47,712	2.22
Yale University	34,241	2.21
UC Los Angeles (UCLA)	37,742	2.21
University of Chicago	16,265	2.13

Organization, country	pa.eu/dashboard 23 SEP : Net contri-	project
(*med school only)	bution (in €)	participations
INSERM, FR	115.160.351	122
Univ of Oxford, UK	76.643.642	74
LSHTM, UK	74.201.528	26
Erasmus MC*, NL	61.255.042	72
Karolinska Inst*., SE	61.171.462	89
Radboud Univ, NL	57.262.658	52
UCL, UK	55.748.799	63
UMC Utrecht*, NL	53.889.035	50
ICL, UK	50.417.535	43
KCL, UK	49.689.847	49
KU Leuven, BE	45.388.558	68
LUMC*, NL	43.742.800	56

 Tabel kiri:
 Dampak dunia:
 dampak sitasi kelompok publikasi ini dibandingkan dengan dampak dunia (rata-rata dunia adalah 1,00). Publikasi WoS: publikasi penelitian dalam

 gabungan domain ilmu praklinis, klinis, & kesehatan antara tahun 2016-2020 sebagaimana terdapat dalam pangkalan data InCites pada 12 Oktober 2021

 Tabel kanan:
 organisasi tersukses dalam program pendanaan penelitian Eropa Horizon2020 – bertema Kesehatan, Perubahan Demografis, & Kesejahteraan, sebagaimana

 diberi peringkat menurut jumlah euro yang diperoleh sebagaimana terdapat pada dasbor UE pada 23 September 2020. Erasmus MC adalah sekolah kedokteran kontinental

 pertama, karena INSERM Prancis adalah organisasi nasional dan dua organisasi tersukses lainnya adalah organisasi Inggris.

Tujuan program PhD di Erasmus MC adalah membuat Anda menjadi peneliti independen yang dapat menyelesaikan berbagai pertanyaan rumit berdasarkan bukti ilmiah. Lulusannya akan memiliki kompetensi untuk menilai penelitian ilmiah dan telah mengambil langkah penting untuk menjadi akademisi biomedis. Mahasiswa PhD disiapkan secara optimal untuk menjadi staf peneliti (klinis) di masa depan di pusat kedokteran universitas, universitas penelitian, lembaga penelitian, dan/atau mengisi posisi staf dan pengambil kebijakan, seperti manajemen universitas biomedis, rumah sakit, dan organisasi kesehatan lainnya, perusahaan biomedis dan farmasi, kementerian, dan banyak lagi.

Inti filosofi pendidikan kami adalah pelatihan ilmiah yang baik membutuhkan pemelajaran aktif. Artinya, kami mengajar mahasiswa PhD dan magister penelitian dalam kelompok kecil atau bahkan terkadang perseorangan, dan berarti bahwa pengetahuan teoretis dan keahlian praktis diajarkan dengan cara terintegrasi. Oleh karena itu, mahasiswa distimulasi untuk secara aktif menggunakan pengetahuan yang baru mereka peroleh, yang menanamkan pengetahuan mereka dan meningkatkan kualitas penelitian mereka. Titik pertemuan ini adalah penggerak penting untuk meningkatkan multidisiplin dan transdisiplin dalam pendidikan kami di semua tingkat. Mahasiswa belajar dari staf pengajar terbaik di bidang mereka dengan pengalaman internasional dan kelompok penelitian yang berkolaborasi dengan kelompok penelitian (inter)nasional lain.

Program PhD biasanya membutuhkan waktu 4 tahun dan kandidat PhD harus memiliki gelar MSc, MD, atau DVM. Dalam ilmu kesehatan, kandidat akan mengombinasikan studi PhD mereka dengan master spesialisasi ilmu kesehatan. Seorang kandidat harus memiliki skor IELTS 7,0 atau TOEFL 100, tetapi saat menempuh pendidikan PhD, keahlian menulis dan presentasi mereka dalam bahasa Inggris lebih disempurnakan.

Pelatihan dan pembimbingan: Sebagai mahasiswa PhD, Anda akan masuk ke Erasmus MC Graduate School yang menawarkan mata kuliah umum dan mata kuliah yang sangat terspesialisasi. Tetapi, program PhD sangat individual dan dalam beberapa bulan pertama, bersama dengan pembimbing Anda, Anda akan mengembangkan program sendiri yang paling sesuai dengan kebutuhan ilmiah Anda dan jalur karier yang Anda inginkan. Hal yang penting adalah kami juga berharap Anda dapat bekerja secara mandiri (kami melatih Anda untuk melakukannya) serta berani mengambil inisiatif dan kami akan menstimulasi Anda untuk bersaing mendapatkan tunjangan perjalanan, penghargaan presentasi terbaik, atau melakukan kegiatan terkait lainnya yang masih merupakan kegiatan ekstrakurikuler.

- Anda akan melakukan penelitian ilmiah mandiri dan mempresentasikan hasilnya dalam tesis.
- Anda akan dibimbing oleh profesor senior (promotor) dan dibantu oleh satu atau dua pembimbing pendamping
- Anda akan mengikuti mata kuliah, seminar, dan konferensi untuk mendapatkan setidaknya 30 poin EC (Anda dapat memilih dari 150 mata kuliah di Sekolah ini dan Anda diizinkan untuk mengikuti mata kuliah di luar Erasmus MC)
- Anda akan berpartisipasi dalam lingkungan penelitian multidisipliner dan multinasional yang canggih dan digerakkan oleh dana hibah
- Tergantung pada proyek Anda, Anda mungkin dapat pergi ke luar negeri (kunjungan penelitian) untuk belajar di lingkungan lain

Tesis PhD Anda: setiap proyek penelitian berbeda, setiap mahasiswa PhD berbeda, dan pengetahuan serta pengalaman lab berbeda juga, karena mahasiswa PhD berasal dari berbagai universitas. Tetapi, kami bangga memiliki salah satu persyaratan ujian PhD tertinggi di dunia. Hal ini memberi Anda keuntungan signifikan saat mengambil langkah berikutnya dalam karier Anda. Untuk mendapatkan bayangan tentang hasil Anda setelah Anda menerima gelar PhD, lihat tabel di bawah ini:

Keluaran 10 mahasiswa PhD Indonesia terakhir pada saat kelulusan PhD mereka di Erasmus MC

University of Origin	no of publications + field specific ranking of the journal of publication	courses & conferences followed abroad	honors & awards obtained during PhD + extracurricular activities
UI, Jakarta	6x top 25%, 2 other	2 courses, 2 conferences	DIKTI + supervising 6 interns, 1 clinical fellow, tutor at clinical lab course
JI, Jakarta	4x top 25%, 3x other publications, 2x under review, 1 book, 1 book chapter	6 conferences	LPDP, teaching at FKUI, teaching at Indonesische Stichting Rotterdam
JI, Jakarta	1x top3, 3x top 25% 4 other publ, 4x in preparation, 1x submitted, 1x under review, 1 book chapter	3 conferences	DIKTI+3; teaching assistant + supervising master students, guest assoc editor Frontiers Microbiology
JnDip, Semarang	1x top 3, 4x top 5, 3x top 10, 4x top 25%, 3x Top 50% journals	1 course, 4 conferences	NWO+4
JnDip, Semarang	2x top 50%, 1x top 75% journals + 3x in preparation	2 courses, 6 conferences	DIKTI + 2
JGM, Yogyakarta	2x top 3,1x top 5, 2x top 10, 5x top 25%, 2x top 50%, 4 top 75% journals	1 course, 1 conference	LPDP + 3
JnPad, Bandung	1x top 5, 1x top 25%, 3x top 50% journals + 3 in preparation	7 conferences	DIKTI + founding board member QOLMARI Indonesia
JI, Jakarta	2x top 10, 1x top 25%, 2x top 50% journals	2 courses, 7 conferences	DIKTI
JB, Malang	1x top 25%, 4x top 50% journals + 1 in preparation	1 conference	DIKTI + teaching (1450 hours)
UnPad, Bandung	2x top 10, 3x top 25%, 4x top 50%, 1x top 75% journals	3 conferences	NWO + 3

Legenda: <u>universitas asal</u> – universitas tempat lulusan PhD memperoleh gelar MD atau MSc, <u>jumlah publikasi + peringkat khusus bidang yang dimiliki jurnal publikasi tersebut</u> – jumlah publikasi lulusan pada waktu sidang tesis PhD, kualitasnya ditunjukkan dengan peringkat jurnal di bidang penelitian mahasiswa PhD tersebut, <u>mata kuliah &</u> konferensi yang diikuti di luar negeri – jumlah konferensi, mata kuliah, dan kunjungan penelitian di luar Belanda dan di luar Indonesia, <u>pengakuan & penghargaan yang</u> <u>diperoleh selama PhD + kegiatan ekstrakurikuler</u> – jumlah hibah & penghargaan, beasiswa atau tunjangan perjalanan, keanggotaan komite atau dewan yang diperoleh, dan kegiatan mengajar.

Setelah tesis Anda hubungan Anda dengan kami tidak akan terputus setelah Anda memperoleh gelar PhD Anda: mengenal staf dan penelitian kami serta memahami dinamika dana hibah penelitian di negara Barat, Anda akan berubah dari mahasiswa pascasarjana menjadi kolega dan mitra penelitian yang penting di luar negeri: banyak dari kolaborasi sukses kami dilakukan dengan alumni kami.

Cara mengajukan aplikasi untuk posisi PhD

Cara menggunakan buklet lowongan ini: Buklet ini adalah gambaran tentang posisi mahasiswa PhD di berbagai lab di berbagai departemen di Erasmus MC. Tetapi, jika Anda menyukai bidang penelitian profesor tertentu yang bukan lowongan PhD profesor tersebut, Anda selalu dapat menghubungi profesor tersebut karena alamat email disebutkan dalam lowongan. Sebagian besar lowongan dibuat secara umum untuk memberi Anda informasi tentang topik yang dipelajarinya, tetapi juga memberi Anda fleksibilitas untuk mengajukan modifikasi pada tema. Anda mungkin juga tidak menemukan jenis penelitian yang Anda minati: buklet ini menunjukkan 50 lowongan mahasiswa PhD, tetapi ada lebih dari 200 profesor senior dan ada 1.500 staf ilmiah. Oleh karena itu, Anda selalu bisa mengunjungi <u>www.erasmusmc.nl</u>, atau <u>laman lowongan</u> dan menghubungi staf Erasmus MC berdasarkan informasi dalam situs web tersebut daripada informasi dalam buklet ini.

Menulis surat motivasi atau surat pengantar: lowongan berisi uraian singkat tentang penelitian dan menunjukkan sedikit publikasi. Ini adalah sumber yang dapat Anda baca lebih lanjut. Pembimbing berharap kandidat PhD menulis <u>surat</u> motivasi yang baik, yang menggambarkan minat mereka pada minat penelitian profesor tersebut dan bagaimana pengalaman kandidat akan cocok dengan atau akan bermanfaat untuk proyek PhD tersebut.

Karena hampir semua mahasiswa PhD di Erasmus MC memiliki posisi mereka berdasarkan dana hibah penelitian atau beasiswa PhD sendiri, disarankan untuk menyebutkan bahwa Anda akan mengajukan aplikasi untuk beasiswa PhD jika diterima oleh profesor tersebut. Ini bisa berupa beasiswa nasional (<u>LPDP</u>, <u>DikBud</u>), beasiswa PhD universitas atau rumah sakit universitas. Memperoleh beasiswa bisa terasa seperti persyaratan, tetapi kami melihatnya sebagai langkah ekstra yang akan berfungsi sebagai bukti kualitas dalam karier Anda di kemudian hari. Ini juga menjadi alasan pembimbing Anda untuk membantu Anda dalam bagian penelitian di aplikasi beasiswa Anda.

Anda diterima oleh seorang profesor, sekarang bagaimana? Setelah Anda menyelesaikan wawancara (atau beberapa wawancara) dan diterima, dalam sebagian besar kasus, Anda akan mengajukan aplikasi beasiswa. Pembimbing Anda akan menawarkan bantuan dalam uraian ilmiah di aplikasi beasiswa PhD Anda dan seringkali Anda membutuhkan Surat Penerimaan untuk aplikasi beasiswa Anda. Pembimbing Anda dapat memperolehnya melalui <u>RDO</u>. Saat Anda mendaftarkan diri untuk beasiswa lain di universitas Anda atau rumah sakit yang berafiliasi dengan universitas, Anda selalu bisa bertanya kepada pembimbing Anda atau menghubungi <u>RDO</u>.

Setelah aplikasi Anda dikirimkan dan, beberapa waktu kemudian, beasiswa Anda diberikan, Anda akan menginformasikannya kepada pembimbing Anda. Pembimbing Anda akan menginformasikannya kepada Departemen Personalia dan Sumber Daya Manusia (SDM) Anda sebagai mahasiswa PhD barunya, dan Anda dapat dihubungi juga oleh staf Erasmus MC lain. Biasanya, Personalia akan menghubungi Anda dua bulan sebelum perkiraan kedatangan Anda.

Dokumen yang Anda perlukan untuk Personalia untuk menyiapkan aplikasi & pendaftaran Anda

- Salinan paspor berwarna (semua halaman yang berisi tulisan dan berstempel);
- bukti asuransi kesehatan dengan pertanggungan di Belanda; Jika belum memilikinya, Anda dapat membuat asuransi kesehatan setelah tiba di Belanda;
- **Bukti kemandirian**: contohnya gaji, hibah, sponsor, pembayaran berkala, surat pengangkatan, atau kontrak kerja.
- Salinan sertifikat yang membuktikan bahwa Anda memiliki kualifikasi yang sesuai untuk melakukan penelitian;
 ijazah Anda. Ijazah harus divalidasi oleh notaris atau pejabat kota yang berwenang;
- Salinan proposal penelitian, yang ditandatangani oleh pembimbing Anda.

Selain dokumen wajib di atas, disarankan agar Anda menyampaikan juga

salinan akta lahir, yang telah dilegalisasi atau dicap apostille untuk memastikan detail pribadi untuk Pangkalan
 Data Catatan Pribadi Kota (GBA).

<u>NB</u> Semua dokumen tersebut harus diterjemahkan ke bahasa Inggris, Belanda, atau Prancis oleh penerjemah resmi.

Department of Biochemistry

Work environment:

Erasmus MC is an internationally recognized centre for highly rated transfer of knowledge and high-quality knowledge development in the fields of illness and health. The research groups at the department of Biochemistry are interested in the understanding of the mechanisms of gene expression control during development and disease.

<u>Peter Verrijzer</u>'s lab aims to understand the mechanisms of gene regulation that underpin development and disease. We are particularly interested in the role of chromatin remodelers in human disease and the coupling between cellular metabolism and epigenetics. We use an integrated approach, combining biochemistry, proteomics, developmental genetics and cell biology. Taking advantage of evolutionary conservation, key regulators are studied both in human cells and in the genetically tractable fruit fly.

Tokameh Mahmoudi's lab aims to translate basic molecular advances in the HIV and HBV field into development and testing of novel therapeutics in the clinic. We delineate the molecular events that lead to HIV latency and HBV– mediated liver tumorigenesis. Parallel projects use unbiased and candidate approaches to identify molecular targets or therapeutic molecules in HIV latency reversal, which we characterize in in vitro latency models and T cells obtained from HIV infected patient volunteers. We also use the human liver organoid technology to model HBV infection and study mechanisms of HBV-induced liver tumorigenesis.

Jeroen Demmers's lab develops mass spectrometry-based methodologies for qualitative and quantitative proteomics analysis. Our research focuses on the analysis of protein post-translational modifications, protein-protein interactions, protein complex composition and analysis of proteome dynamics. The ultimate goal is to develop analytical tools to better understand how cellular processes are controlled at the molecular level in health and disease.

Selected publications:

<u>Verrijzer</u>

2017 Mohd-Sarip A et al **Cell Reports** 2014 Reddy BA et al **Molecular Cell** 2013 Moshkin YM et al **PLoS Genet** 2012 Mohd-Sarip A et al **Science**

Mahmoudi

2018 Marian C et al **Cell Chem Biol** 2018 Palstra R-J et al **Science Advances** 2016 Stoszko M et al **EBioMedicine** 2012 Li V et al **Cell**

Demmers:

2017 Sap KA et al **J Proteome Res** 2016 Urbán N et al **Science** 2016 Yu N et al **Curr Biol** 2012 Schwertman et al.**Nat. Genet**

Qualifications and skills:

We are looking for highly motivated PhD students that have received excellent scientific and practical training in the areas of Molecular Virology, Molecular Biology, Proteomics, or bioinformatics to join our research teams. The Biochemistry department has a modern infrastructure and facilities. We have in house access to the very efficient and up-to-date core proteomics, genomics, and bioinformatics and in house high through put DNA and RNA sequencing facilities. We have an MLII facility for HBV work and have access to and use the MLIII and MLII (biosafety level 2 and 3) and MLI cell culture facilities.

We offer: High quality state-of-the-art project, supervision, lab facilities and infrastructure, and training. We will cover Laboratory costs. Your salary and living expenses will be covered by your University or Scholarship Council.

Department of Biochemistry

Combination

School/Department:	Department of Biochemistry and Department of Pathology, Erasmus MC		
Supervisor	Prof. dr. Tokameh Mahmoudi, PhD, <u>t.mahmoudi@erasmusmc.nl</u>		
information:	Lab webpage: <u>Mahmoudilab.com</u>		
Selected grants: ERC StG, Health Holland, ZonMW 2019		W 2019	
	Selected publications:		
	2021 Nature Communications 12(1):2475	2020 Science Advances 6(32):6617-6629	
	2020 Journal of Virological Methods.	2020 Viruses. 12(9):E973.	
	2019 Current Opinion in Virology.	2019 Pharmacol Res. 2019 Jan;139:524-534.	
	2020 bioRxiv	2018 Cell Chemical Biology 25(12):1443-1455.e14.	
	2018 Science Advances 4(2):e1701729.	2016 EBioMedicine. 3:108-121.	

Project Title:

Abstract:

antiretroviral therapy effectively halts HIV replication and has significantly reduced AIDS-associated mortality. However, cART is not curative, it has side-effects, and apart from the costs of lifelong therapy, the capable of global roll-out of cART, particularly in resource-limited countries, remains an ongoing HIV, challenge. HIV persists because the integrated provirus can their remain in a nonproductive latent state, defined by the action. absence of HIV-1 gene [3] The expression. Because of this reservoir of latently HIV-1 infected cells, interruption of factors cART leads to a rapid rebound of unrestricted viral replication, necessitating life-long treatment. Ongoing progress in understanding the molecular mechanisms that control HIV transcription and latency has led to the development of strategies to target the reservoir, to stimulate the virus to emerge out of latency, coupled to either induction of death in the infected reactivated cell or its immune

clearance. World no 24 in Infectious Diseases World no 30 Biomedical Sciences

Requirements of candidate:

HIV Cure: mechanisms, drug discovery, clinical study and valorization

We use various cell based and patient-derived models of HIV latency to screen for, identify, characterize, and clinically translate potential novel therapeutics toward HIV cure:

[1] An innovative approach to eliminate HIV-1-infected cells emerging out of latency is to pharmacologically reactivate viral expression and concomitantly trigger intracellular pro-apoptotic pathways in order to selectively induce cell death (ICD) of infected cells.

[2] Using a medium through-put screen of fungal metabolites combined with HIV latency reversal bioassays and state of the art fractionation coupled to MS and NMR bioassays, we identify molecules

activating latent characterize mechanisms of unbiased identification of physically associated with the latent HIV-1 provirus would



be highly valuable to unravel the molecular correlates of latency and develop new latency reversal agents. But, due to technical limitations, this has not been possible.

We developed dCas9 targeted chromatin and histone enrichment strategy coupled to mass spectrometry (Catchet-MS) to isolate the latent HIV-1 promoter and identified novel and previously known factors physically associated with potentially repressing the latent LTR, and will investigate the molecular mechanisms by which they function. For one of the candidates bound, we found the FDA approved IKZF1 targeting thalidomide analogues reversed latency in CD4+T-cells isolated from virally suppressed HIV-1 infected participants.

[4] We identified the BAF complex as a central player in repressing HIV transcription, highlighting it as a potential target to reverse HIV latency. In collaboration we found that small-molecule inhibition of BAF re-activates latent HIV in a spectrum of primary models as well as in cells obtained from HIV-infected patients using drug screens. We also found macrolactam scaffold BAF inhibitors to be potentially potent latency reversal agents.

•We are looking for a highly motivated PhD student who has received excellent scientific and practical training in the areas of Molecular Virology or Molecular Biology who also has some basic training or interest in bioinformatics to join our research team.
•The student should be fluent in English (<i>English speaking countries & Netherlands:</i> no requirement; Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs).
•We offer: Supervision, lab facilities and infrastructure, and training. We will cover Laboratory costs.
 As a candidate PhD student at Erasmus MC, your salary and living expenses will be covered by your University or Scholarship Council.

Department of Biochemistry

School/Department:	Department of Pathology and Department of Biochemistry, Erasmus MC	
Supervisor	Prof. dr. Tokameh Mahmoudi, PhD, <u>t.mahmoudi@erasmusmc.nl</u>	
information:	Selected grants: ERC StG, Health Holland, ZonMW 2019	
	Selected publications (* as last author):	
	2021 Elife 10:e60747. Application of human liver organoids as a patient-derived primary model for HBV	
	infection and related hepatocellular carcinoma*	
	2021 Nature Communications. doi: 10.1038/s41467-021-22608-z. Selective cell death in HIV-1-infected	
	cells by DDX3 inhibitors leads to depletion of the inducible reservoir*	
	2021 Cell Death Dis. 12(7):641. Clinical stage drugs targeting inhibitor of apoptosis proteins purge	
	episomal Hepatitis B viral genome in preclinical models.	
	2021 Cancer Lett. 506:35-44. 3D human liver organoids: An in vitro platform to investigate HBV infection, replication and liver tumorigenesis*	
	<u>2012 Cell</u> 149(6):1245-56. Wht pathway activation through inhibition of proteosomal bcatenin	
	degradation within the intact endogenous Axin1 complex*	
Project Title:	Human liver organoid-tumoroid platform in study of HBV infection and	
	tumorigenesis	
Main methodology and	Abstract: Persistent Hepatitis B virus (HBV) infection remains the leading cause of liver	
techniques 3D liver	cirrhosis and hepatocellular carcinoma world-wide. However, the molecular events that occur	
organoid cultures from	as consequence of HBV infection and which mediate onset of hepatocellular carcinoma have	
healthy donor, HBV infected	remained elusive because of lack of a relevant primary untransformed model system. My	
and hepatocellular	group, in collaboration with the HUB has recently developed a patient-derived HBV infected	
carcinoma patients, Next	human liver organoid model system (de Crignis 2021), using the adult stem cell human liver	
generation sequencing	organoid/tumoroid technology (Huch 2015), which allows long term culturing and analysis of	
analysis of chromatin and	HBV infected patient or healthy donor livers providing a platform suitable for antiviral drug	
gene expression (ChIP-seq	screening and examination of HBV-induced mechanisms of liver pathogenesis and HCC.	
and RNA-seq) , High	Human liver organoids are infected with both recombinant virus as well as HBV infected	
resolution imaging	patient serum and determinants of infection and viral replication are examined. We generate	
(confocal, fluorescence	transgenic organoids to study the function of viral and host factors and perform drug and	
microscopy), Flow	toxicity screens using the HBV liver organoid platform and examine the role of various	
Cytometry Activated Cell	pathways implicated in liver cancer such as Wnt-bcatenin (Li VS 2012), and epigenetic	
Sorting, Lentiviral	regulators.	
transduction and gene		
editing, molecular biology		
and molecular virology		
techniques.		
tali aliana		
Lab webpage:		
Mahmoudilab.com		
world no 14		
<u>Gastroenterology &</u>		
Hepatology		
<u></u>		
World no 30 Biomedical		
<u>Sciences</u>		
Requirements of	• We are looking for a highly motivated PhD student who has received excellent scientific and practical training in the areas of Molecular Virology or Molecular Biology who also has some basic training or interest in Bioinformatics to join our research	
candidate:	team.	
	• The student should be fluent in English (English speaking countries & Netherlands: no requirement; Other countries: IELTS 7.0	
	 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs). We offer: Supervision, lab facilities and infrastructure, and training. We will cover Laboratory costs. 	
	 We other: supervision, (ab facilities and intrastructure, and training, we will cover Laboratory costs. As a candidate PhD student at Erasmus MC, your salary and living expenses will be covered by your University or Scholarship 	
	Council.	

Department of Biostatistics

School/Department:	Department of Biostatistics, Erasmus MC
Supervisor information:	Prof. dr. Dimitris Rizopoulos, (promotor, <u>d.rizopoulos@erasmusmc.nl)</u>
	Dr. Joost van Rosmalen (co-promotor, j.vanrosmalen@erasmusmc.nl)
<u>World no 21 Public,</u>	See <u>www.drizopoulos.com</u> and
Environmental & Occupational	https://www.scopus.com/authid/detail.uri?authorId=26041070200 for a personal website and an
Health 2021	overview of publications. The most relevant publications on this topic are:
	-J. van Rosmalen, D. Dejardin, Y. van Norden, B. Löwenberg, E. Lesaffre (2017). Including historical data in the
	analysis of clinical trials: Is it worth the effort? Statistical Methods in Medical Research.
	-Hatswell A, Freemantle N, Baio G, Lesaffre E, van Rosmalen J (2020). Summarising salient information on
	historical controls: A structured assessment of validity and comparability across studies. Clin Trials. -Banbeta A, van Rosmalen J, Dejardin D, Lesaffre E (2018). <i>Modified power prior with multiple historical trials for</i>
	binary endpoints. Stat Med
Project Title:	How to assess the value of historical controls in Bayesian dynamic borrowing methods
Abstract:	Consider the common situation where a clinical trial is planned, say on a new treatment for
	Alzheimer's disease, and data from previous trials are available. The intervention treatment
	tends to differ across trials, but the control treatment often remains the same. We might then
	add the controls of the previous trials to the analysis of the current (newly planned) trial, to
	increase the statistical power and reduce the sample size. However, care must be taken to
	ensure that these historical data are sufficiently comparable to the current study, to avoid a bias
	in the estimates. Several Bayesian statistical methods have been developed that include the
	historical data when it is sufficiently similar to the current data, but downweight or even discard
	the historical data in case of substantial differences. The main methods are the power prior
	(Ibrahim & Chen, Statistical Science 2000), the meta-analytic predictive prior (Neuenschwander
	et al., Clin Trials 2010) and the commensurate prior (Hobbs et al., Bayesian Anal 2012). Despite
	the wide range of available methods, it's not clear which method performs best.
	In this project we will focus on determining which of the available methods is best suited for
	practical use, what settings should be used for that method and on developing a framework with
	appropriate metrics (e.g. power and type I error rate) to compare different methods. The meta-
	analytic predictive prior will be the starting point.
	Research questions include:
	 How should frequentist characteristics of borrowing methods be assessed? What is the best upputs make borrowing methods rebust against prior data conflict?
	- What is the best way to make borrowing methods robust against prior-data conflict?
	- How should we choose the settings (e.g. the prior) of these dynamic borrowing methods
	to optimize the tradeoff between power and type I error rate?
	- How can we justify the choice for a borrowing method based on what we know about
	the similarity of the historical and the current data?
	These borrowing methods will be applied to real-life case studies (e.g. we have a case study on a
	series of trials for Alzheimer's disease) and simulated data.
	Keywords: Bayesian statistics, biostatistics, historical data, power prior, meta-analytic predictive
	prior
Requirements of candidate:	We're looking for an enthusiastic student with a background (master's degree) in biostatistics or statistics who is interacted in developing and applying new biostatistical mathedology. Knowledge of Bayesian statistics is a
	interested in developing and applying new biostatistical methodology. Knowledge of Bayesian statistics is a prerequisite. A good command of the English language (especially writing) is also necessary.
	ירבי בקעומונים. א צטטע נטווווזמווע טו נוופ בווצוומו ומווצעמצים (פאפטומווץ שרונווצ) וא מוגט וופניפאמו א.
	We offer a good working environment with a friendly atmosphere and constructive scientific supervision in the
	Department of Biostatistics of Erasmus MC, Rotterdam, the Netherlands. The department is well known for its
	expertise on methods for analyzing longitudinal data (joint modeling and other methods), Bayesian statistics and
	the analysis of historical data. In addition to the project outlined above, we can also facilitate PhD projects on
	other topics.
	The scholarship will, at least, cover subsistence allowance and an international airplane ticket. We're able to
	provide help with the scientific part of your scholarship proposal.
	English language requirement: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Cardiology, section electrophysiology, Erasmus MC
Supervisor	•Prof dr. Natasja MS de Groot
information:	•Email: <u>n.m.s.degroot@erasmusmc.nl</u>
-	•Website: <u>https://www.linkedin.com/in/prof-dr-natasja-de-groot-md-phd-emc-65760662/</u>
World no 23 in Cardiac &	https://www.erasmusmc.nl/en/research/researchers/groot-natasja-de ,
Cardiovascular Systems	https://www.medicaldelta.nl/onderzoek/medical-delta-cardiac-arrhythmia-lab
<u>caratovascular systems</u>	•Grants: EU-LSH, Dutch-German Heart Foundation grant, Cardiovascular research Netherlands, personal grants: Dutch
	Heart Foundation Junior Staffmember, VIDI; multiple companies (e.g. Johnson&Johnson, Bayer) Most important
	publications: Zhang, D., et al. (2019) Nature Communications, Calkins, H., Heart Rhythm, de Groot, N.,
	(2016) Circulation-Arrhythmia and Electrophysiology; Knol, W. G., et al. (2019). Heart Rhythm,
Drainat Titla	Starreveld, R., (2019) Europace, Kharbanda R. (2020) JACC EP.
Project Title:	Innovation in Diagnosis and Therapy of Cardiac Arrhythmias
Abstract:	Our projects are aimed at unravelling the pathophysiology of complex cardiac tachyarrhythmias, developing and testing developing novel diagnostic tools (in close collaboration with Technical university Delft) and therapies for cardiac arrhythmias. Main topics are high resolution mapping studies of cardiac arrhythmias in particular atrial fibrillation, unravelling bio-electrical mechanisms
	of (post-operative) cardiac arrhythmias, dysrhythmias in patients with congenital heart disease and neuromodulation of atrial fibrillation. For this purpose, we have developed a unique way of recording and processing cardiac signals to perform mapping procedures in the surgical rooms and catheterization laboratory. In addition, we have access to biomimetic set ups for tissue slices and
	an ex-vivo-heart perfusion model.
	Our innovative scientific contributions include: discovery of novel mechanisms underlying persistence of atrial fibrillation, introduction endovascular mapping approach guiding ablative therapy of atrial tachyarrhythmias in patients with congenital heart disease, development of a novel, intra- operative epicardial mapping approach, discovery of the role of Bachmann's bundle in development of atrial tachyarrhythmias, performed worldwide the first high resolution mapping studies in pediatric patients, discovery conduction properties in pediatric patients with congenital
	heart disease.
	In our cardiac bio-electricity lab, we combine expertise of developmental biology, cardiac electrophysiology with macro- and microscopic cardiac morphology. We perform clinical and experimental studies in surgical rooms, EP labs, outpatient clinic and animal lab. We have several multi-disciplinary collaborations and electrical-, biomechanical engineers, a variety of medical doctors and molecular biologist are part of our research group.
	Keywords: cardiac surgery, electrophysiology laboratory, biomarkers, human-, animal-, clinical-,
	experimental mapping studies, electrical activity, ECG analysis, electrograms, biomarkers and medical
	technology.
Requirements of	• We are looking for highly motivated, hardworking students to join our very international team. Our strength is
candidate:	in using team work to tackle large scientific questions.
	 Master degree or MD Scholarship that will, at least, cover subsistence allowance and international airplane ticket (we could help
	with the scientific part of your scholarship proposal)
	English language requirement:
	 English speaking countries & Netherlands: no requirement
	• Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Cardiology, Erasmus MC
Supervisor information:	Dr. HMM van Beusekom, Dr. Majoor-Krakauer, Dr. IJpma, Dr. Vreeken
	Email: h.vanbeusekom@erasmusmc.nl
World no 23 in Cardiac &	Website: Department - Cardiology (erasmusmc.nl)
Cardiovascular Systems	Grants: 2020-2024 Private Foundation: Aortic Aneurysm disease
	 2020-2024 Private Poundation. About Aneurysin disease 2018-2022 ZonMW Coronary stent in a box and on a chip
	 2016-2022 CVON CONTRAST Development of gyrencephalic stroke models, thrombus biobank analyses
	 2014-2018 ZonMW Imaging drug and scaffold metabolomics in coronary artery disease
	• 2013 Thrombosis foundation <u>Functional three-dimensional architecture of the coronary thrombus</u>
	Most important publications:
	- Consensus standards for acquisition, measurement, and reporting of intravascular OCT GJ Tearney, E Regar, T Akasaka, et al, Journal of the American College of Cardiology 59 (12), 1058-1072; 2012
	 Marked inflammatory sequelae to implantation of biodegradable and nonbiodegradable polymers in porcine coronary
	arteries WJ Van der Giessen, AM Lincoff, RS Schwartz, HMM Van Beusekom, et al, Circulation 94 (7), 1690-1697; 1996
	- Endothelial progenitor cell capture by stents coated with antibody against CD34First In Man J Aoki, PW Serruys, H van
	 Beusekom, et al, Journal of the American College of Cardiology 45 (10), 1574-1579; 2005 Intracoronary optical coherence tomography and histology at 1 month and 2, 3, and 4 years after implantation of
	everolimus-eluting bioresorbable vascular scaffolds in a porcine Y Onuma, PW Serruys, LEL Perkins, et al, Circulation 122
	(22), 2288-2300; 2010
	 Reduction in thrombotic events with heparin-coated Palmaz-Schatz stents in normal porcine coronary arteries. PA Hårdhammar, HMM van Beusekom, HU Emanuelsson, et al, Circulation 93 (3), 423-430; 1996
	 Mutations in SMAD3 cause a syndromic form of aortic aneurysms and dissections with early-onset osteoarthritis. van de
	Laar IM, Oldenburg RA, Pals G. et al. Nat Genet. 2011;43(2):121-6
	- Cardiac Phenotypes, Genetics, and Risks in Familial Noncompaction Cardiomyopathy. J.I. van Waning, K. Caliskan, M.
	Michels et al. J Am Coll Cardiol 2019;73 (13);1601-11
Project Title:	Human disease model technology and mathematical modelling for arterial
-	interventions in coronary arteries and aortic aneurysms
Abstract:	This line of investigation is a collaboration between several Erasmus MC departments (Clinical
	genetics (Majoor-Krakauer), Pathology (IJpma), Cardiology (van Beusekom, Vreeken) and Delft
	University of Technology (van Steijn). Our group aims to develop animal free models to study
	vascular disease and improve treatment strategies. In particular, we focus on coronary
	interventions and aortic aneurysms.
	Coronary interventions. In this project we culture coronary arteries in a bioreactor (VABIO),
	which allows real-time ultrasound and OCT imaging to study coronary atherosclerosis and
	vascular responses to percutaneous coronary interventions (PCI) especially drug eluting
	stents. We specifically study drug distribution in the arterial wall and how this relates to
	vascular disease. To that end we also develop organ-on-a-chip (OOC) approaches in
	collaboration with the Delft University of Technology (TUD).
	Aortic aneurysms. This project aims to develop human disease models to mimic and predict
	aortic aneurysm formation. This will help to reveal potential risks for AA disease development
	as well as predicting outcome after treatment using endovascular repair strategies (EVAR) on
	the aortic wall.
	PhD positions would be possible in the
	1. Bioreactor culture arena for coronary arteries and aortae, and the development of OOC
	approaches for PCI and EVAR.
	2. A technology-oriented PhD position that deals with modelling of cellular and chemical
	processes in the arterial wall in collaboration with TUD.
Requirements of	• We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using
candidate:	teamwork to tackle large scientific questions and thus require a student with good communication skills.
	 Master degree or MD Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific
	part of your scholarship proposal)
	English language requirement:
	 English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Cardiology Erasmus MC
Supervisor information:	Dr. HMM van Beusekom and Dr. J Bobii Gibert
	Email: <u>h.vanbeusekom@erasmusmc.nl</u> or j.bobiigibert@erasmusmc.nl
World no 23 in Cardiac &	Website: Department - Cardiology (erasmusmc.nl)
Cardiovascular Systems	Grants:
Caralovascular Systems	2020-2024 Private Foundation: Aortic Aneurysm disease
	2020-2022 Erasmus MC grant: Human disease model technology
	2018-2022 ZonMW <u>Coronary stent in a box and on a chip</u>
	2016-2023 <u>CVON CONTRAST</u> Development of gyrencephalic stroke models, thrombus biobank analyses
	2014-2018 ZonMW Imaging drug and scaffold metabolomics in coronary artery disease
	2013 Thrombosis foundation <u>Functional three-dimensional architecture of the coronary thrombus</u>
	Most important publications:
	 Mechanical Characterization of Thrombi Retrieved With Endovascular Thrombectomy in Patients With Acute Ischemic Stroke. Boodt N, Snouckaert van Schauburg PRW, Hund HM et al Stroke. 2021 Aug;52(8):2510-2517. doi: 10.1161/STROKEAHA.120.033527. PMID: 34078112 Endovascular treatment for calcified cerebral emboli in patients with acute ischemic stroke. Bruggeman AAE, Kappelhof M, Arrarte Terreros N, et al; MR CLEAN Registry Investigators. J Neurosurg. 2021 Apr 2:1-11. doi: 10.3171/2020.9.JNS201798. Consensus standards for acquisition, measurement, and reporting of intravascular optical coherence tomography studies: a report from the International Working Group for GJ Tearney, E Regar, T Akasaka, et al, Journal of the American College of Cardiology 59 (12), 1058-1072; 2012 Marked inflammatory sequelae to implantation of biodegradable and nonbiodegradable polymers in porcine coronary arteries WJ Van der Giessen, AM Lincoff, RS Schwartz, HMM Van Beusekom, et al, Circulation 94 (7), 1690-1697; 1996 Endothelial progenitor cell capture by stents coated with antibody against CD34: the HEALING-FIM (Healthy Endothelial Accelerated Lining Inhibits Neointimal Growth-First In Man J Aoki, PW Serruys, H van Beusekom, et al, Journal of the American College of Cardiology 45 (10), 1574-1579; 2005 Intracoronary optical coherence tomography and histology at 1 month and 2, 3, and 4 years after implantation of everolimus-eluting bioresorbable vascular scaffolds in a porcine Y Onuma, PW Serruys, LEL Perkins, T Okamura, N Gonzalo, et al, Circulation 12 (22), 2288-2300; 2010 Reduction in thrombotic events with heparin-coated Palmaz-Schatz stents in normal porcine coronary arteries. PA Hårdhammar, HMM van Beusekom, HU Emanuelsson, et al, Circulation 93 (3), 423-430; 1996 Long-term endothelial dysfunction is more pronounced after stenting than after balloon angioplasty in porcine coronary arteries. HMM van Beusekom, DM Whelan, SH Hofma, et al, Journal of the
Project Title:	Acute ischemic stroke in a large gyrencephalic animal model
Abstract:	In a collaborative project with Erasmus MC departments of Neurology, Radiology and Neurosurgery we developed a swine model of temporary MCA occlusion (clips) to induce focal ischemia-reperfusion and study incomplete microvascular reperfusion and cerebral vasomotor tone. We use imaging techniques <i>to asses reperfusion, cerebral blood flow and infarct size.</i> Histology and (immuno)histochemistry are used to further characterize infarct size and composition. We are looking for a PhD student to further develop the stroke model using vasomotor tone studies <i>in-vivo,</i> using high resolution imaging techniques, and <i>ex-vivo,</i> using organ bath studies or thin brain slice vasoreactivity assays. We would also welcome a PhD student to further develop cognitive assays for long-term follow-up after acute ischemic stroke.
Requirements of candidate:	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills. Master degree or MD Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) English language requirement: English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Cardiology Erasmus MC
Supervisor information:	Dr. HMM van Beusekom, Dr. J. BobiiGibert
, ,	• Email: h.vanbeusekom@erasmusmc.nl or j.bobiigibert@erasmusmc.nl
World no 23 in Cardiac &	Website: Department - Cardiology (erasmusmc.nl)
Cardiovascular Systems	Grants:
Cardiovascular Systems	2020-2024 Private Foundation: Aortic Aneurysm disease
	2020-2022 Erasmus MC grant: Human disease model technology
	• 2018-2022 ZonMW <u>Coronary stent in a box and on a chip</u>
	• 2016-2023 <u>CVON CONTRAST</u> Development of gyrencephalic stroke models, thrombus biobank analyses
	2014-2018 ZonMW Imaging drug and scaffold metabolomics in coronary artery disease
	• 2013 Thrombosis foundation <u>Functional three-dimensional architecture of the coronary thrombus</u>
	Most important publications:
	 Mechanical Characterization of Thrombi Retrieved With Endovascular Thrombectomy in Patients With Acute Ischemic Stroke. Boodt N, Snouckaert van Schauburg PRW, Hund HM et al Stroke. 2021 Aug;52(8):2510-2517. doi: 10.1161/STROKEAHA.120.033527. PMID: 34078112
	 Endovascular treatment for calcified cerebral emboli in patients with acute ischemic stroke. Bruggeman AAE, Kappelhof M,
	Arrarte Terreros N, et al; MR CLEAN Registry Investigators. J Neurosurg. 2021 Apr 2:1-11. doi: 10.3171/2020.9.JNS201798.
	• Consensus standards for acquisition, measurement, and reporting of intravascular optical coherence tomography studies: a report from the International Working Group for GJ Tearney, E Regar, T Akasaka, et al, Journal of the American College of
	Cardiology 59 (12), 1058-1072; 2012
	 Marked inflammatory sequelae to implantation of biodegradable and nonbiodegradable polymers in porcine coronary arteries WJ Van der Giessen, AM Lincoff, RS Schwartz, HMM Van Beusekom, et al, Circulation 94 (7), 1690-1697; 1996
	 Endothelial progenitor cell capture by stents coated with antibody against CD34: the HEALING-FIM (Healthy Endothelial
	Accelerated Lining Inhibits Neointimal Growth-First In Man J Aoki, PW Serruys, H van Beusekom, et al, Journal of the
	American College of Cardiology 45 (10), 1574-1579; 2005
	 Intracoronary optical coherence tomography and histology at 1 month and 2, 3, and 4 years after implantation of everolimus-eluting bioresorbable vascular scaffolds in a porcine Y Onuma, PW Serruys, LEL Perkins, T Okamura, N Gonzalo, et al, Circulation 122 (22), 2288-2300; 2010
	 Reduction in thrombotic events with heparin-coated Palmaz-Schatz stents in normal porcine coronary arteries. PA
	Hårdhammar, HMM van Beusekom, HU Emanuelsson, et al, Circulation 93 (3), 423-430; 1996
	• Long-term endothelial dysfunction is more pronounced after stenting than after balloon angioplasty in porcine coronary arteries. HMM van Beusekom, DM Whelan, SH Hofma, et al, Journal of the American College of Cardiology 32 (4), 1109-1117; 1998
Project Title:	Arterial thrombosis in acute myocardial infarction and acute ischemic stroke
Abstract:	We have a biobank of coronary thrombi aspirated from patients suffering an acute coronary
	syndrome containing thrombi and periprocedural plasma and contains thrombus and plasma
	samples of more than 900 patients. We want to investigate the relation between thrombus
	composition, plasma biomarkers and patient outcome.
	We aim to do the same as host of the Dutch biobank and core lab for thrombi collected during
	stroke treatment in the MRCLEAN studies. This growing biobank now contains over 2000
	sample and is connected to clinical databanks (radiology and neurology).
	This line of investigation is a collaboration between the departments of Cardiology,
	Neurology, Radiology and Pulmonary Disease at Erasmus MC. We study the relation between
	thrombus composition and clinical data such as etiology of thrombosis, patient outcome and
	imaging data.
Requirements of	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to teakle large scientific questions and thus requires a student with good communication skills.
candidate:	 using team work to tackle large scientific questions and thus requires a student with good communication skills. Master degree or MD
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the
	scientific part of your scholarship proposal)
	English language requirement:
	• English speaking countries & Netherlands: no requirement
	 Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Epidemiology	Department of Cardiology, Erasmus MC
Supervisors information:	• Dr. Maryam Kavousi, MD, PhD	Professor Dirk J.G.M Duncker, MD, PhD
. ,	Email: <u>m.kavousi@erasmusmc.nl</u>	Email: <u>d.duncker@erasmusmc.nl</u>
World no 21 Public,	Website:	Website: <u>https://www.erasmusmc.nl/en/</u>
Environmental &	https://www.erasmusmc.nl/en/research/groups	research/departments/cardiology
	/cardiometabolic-epidemiology	Grants and Awards:
Occupational Health 2020	Grants and Awards:	NATO Science Fellowship (1991)
	AXA Research Fund (2012)	American Heart Association (1992, 1994)
World no 23 in Cardiac &	• <i>IDF</i> (2014)	Royal Dutch Academy of Sci. Fellowship (1995)
Cardiovascular Systems	 Prestigious UNESCO-Loreal Fellowship 'For Women in Science' (2014) 	 Dutch Heart Foundation (1999, 2007) Prestigious Dutch Heart Foundation Established Investigator
	 Prestigious ZonMw VENI Grant (2015) 	Fellowship (2000)
	 COLCIENCIAS (2016) 	• Erasmus MC Grant (2008)
	• Erasmus MC Mrace Grant (2016, 2019)	• European Space Agency Grant (2004)
	Netherlands Organisation for Scientific Research	• US Navy Grant (2007)
	(2017, 2017, 2019, 2020, 2020)	Center for Translational Mol. Med. Grant (2008)
	• Dutch Heart Foundation (2017, 2019, 2020)	• EU-FP7-Health-2010 Grant (2010)
	NIH (2019, 2020) Surgeon Commission Marizon 2020 (2020)	 Dutch CV Research Grants (2012, 2014, 2017) Wellcome Trust Grant (2017)
	 European Commission Horizon 2020 (2020) European Commission Horizon 2020 – Innovative 	 Wencome Trast Grant (2017) Prestigious Gabor Kaley Award from the American Physiological
	Medicines Initiative (IMI) (2020)	Society and the Microcirculatory Society (2020)
	European Society of Cardiology Viviane Conraads	Most important publications:
	Outstanding Achievement Award (2020)	• Circ Res 2007;100:1079-88 / 2008;102:795-803
	Young Academy of The Royal Netherlands Academy	• Physiol Rev 2008;88:1009-86
	of Arts and Sciences (2020) Dutch Cardiovascular Alliance (2020) 	• Circ Heart Fail 2009;2:233-42 / 2016;18:588-98
	 Dutch Cardiovascular Alliance (2020) Most important publications: 	 Circulation 2012;126:468-78 Comprehensive Physioly 2012:2:321-447
	 BMC Medicine 2020; 18:263. 	 Comprehensive Physioly 2012;2:321-447 JACC Cardiovasc Interv 2015;8:1990-99
	 Heart 2020; 1062:133-9. / 2019;105:1414-22. 	 Basic Res Cardiol 2016;111:61 / 2020:115:21
	• Lancet 2019;394:2173-83.	• Cardiovasc Res 2018;114:954-64.
	• Circulation 2019;139:e1019-20.	• Cardiovasc Res 2020;116:741-755 / 756-770
	• JACC 2019;74:1420-21.	• Eur Heart J 2015;36:3134-46 / 2017;38:1951-58
	• Diabetologia 2019;62:1581-90.	• Eur Heart J 2020;41:1687-96 / 2020 (PMID32626906)
	• Circulation Research 2017 121:1392-400	Eur J Heart Fail 2018;20:89-96 Braumweld's Heart Disease 11th Ed. 2018. Ch E7
	 JAMA Cardiology 2017 2:986-94. JAMA 2016 316:2126-34. / 2014 311:1416-23. 	 Braunwald's Heart Disease 11th Ed, 2018, Ch 57 ESC Textbook of Sports Cardiol 2019 Ch 1.2.4
	 JAMA 2010 510:2120-54: 7 2014 511:1410-25: JAMA Cardiology 2016 1:767-76. 	
Project Title:		rdiovascular changes in women and men
Abstract:	Heart failure is largely known as a disease of	the elderly. It has turned out as a global pandemic
		e and is increasing in prevalence. Heart failure is
		ortality, despite advances in medical therapy. Aging
	-	protective mechanisms and growing disease processes
	5 5	ure. This project outlines the link between (normal)
	•	
		of cardiovascular function and development of heart
	•	croscopic changes in cardiovascular structure and
	-	diseases associated with aging. The project will be
	•	artments of Experimental Cardiology (Professor Dirk
	Duncker) and Epidemiology (Dr. Maryam Kav	ousi) and will cover the epidemiology, pathophysiology,
	and prognosis of heart failure from basic labc	pratory studies (Experimental Cardiology) to population-
	based studies (Department of Epidemiology).	Due to differences in cardiovascular structure and
		ake a sex-specific approach throughout the project. This
		f ageing process and transition from a healthy heart to
		aid in appropriate and effective primary prevention
	strategies for both women and men.	and in appropriate and effective printary prevention
Poquiromonto of	-	ring student to join our very international team. Our strength is in using
Requirements of	• We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills.	
candidate:		Ils in laboratory molecular techniques and epidemiology
	Scholarship that will, at least, cover subsistence	allowance and international airplane ticket (we could help with the
	scientific part of your scholarship proposal)	
	English language requirement: Statistics and statistics & Nathority	ander no roquiromont
	 English speaking countries & Netherla O Other countries: IELTS 7.0 (min 6.0 fo 	ands: no requirement r all subs), TOEFL 100 (min 20 for all subs)
	• Other countries: IELTS 7.0 (min 6.0 for	1 uli subsj, TOLI E 100 (11111 20 JUL Uli subsj

The Department of Cell Biology performs top level research at the cutting edge of life and biomedical sciences. The department is truly multi-disciplinary, with expertise in -omics and single-cell technologies, perturbation approaches, and advanced imaging. Research is supported by a team of mathematical biologists. While research is mostly of a fundamental nature, the department strives to apply basic knowledge to health care, for example by improving diagnostics and therapies.

The Department of Cell Biology focusses on:

- Line 1. The regulation of gene expression as a means to establish cell type and fate;
- Line 2. The organization of the cell nucleus, with a focus on chromatin folding and remodeling;
- Line 3. Molecular and cell biological studies of the microtubule cytoskeleton.

Realizing that cells are contiguous entities, connecting the research lines is an important departmental effort. For example, nuclear processes can be viewed both as an endpoint of signal transduction cascades emanating from cell fate-determining factors, but also as a starting point of cellular identity; communication between these processes is mandatory and is regulated a.o. by the cytoskeleton. The department focusses on the functions of molecule(s) and molecular networks in hematopoietic and neural stem/progenitor cells, and, more recently, on cardiomyocytes. It studies individual cells, populations, tissues and organs, and animal models and humans.

The Department of Cell Biology has a strong tradition of intra-departmental interactions, and has (international) collaborations with teams from other top institutes and consortia. The department has an excellent reputation in training top quality PhD students; it currently has about 30 PhD students. The senior PIs are Danny Huylebroeck (head of department), Maarten Fornerod, Niels Galjart, Frank Grosveld, Gert Jansen, Sjaak Philipsen, Raymond Poot, Wilfred van IJcken (also associated with the genomics core facility), Derk ten Berge. Junior PIs are Eskeatnaff Mulugeta, Ana Ruiz-Saenz, Ralph Stadhouders (also with Pulmonology), Debbie van den Berg, Tamar van Dijk, and Jeffrey van Haren. Please, see <u>www6.erasmusmc.nl/cellbiology/research/research-groups</u> for a more extensive description of the various research projects and groups in the department.

Five example publications illustrating the research carried out at the department:

Borg J et al. (2010). Haploinsufficiency for the erythroid transcription factor KLF1 causes hereditary persistence of fetal hemoglobin. **Nature Genetics** 42, 801-805.

Quevedo M et al. (2019). Mediator complex interaction partners organize the transcriptional network that defines neural stem cells. **Nat Commun** *10*, 2669.

ten Berge D et al. (2011). Embryonic stem cells require Wnt proteins to prevent differentiation to epiblast stem cells. **Nature Cell Biology** 13, 1070-1075.

Yu N et al. (2016). Isolation of Functional Tubulin Dimers and of Tubulin-Associated Proteins from Mammalian Cells. **Curr Biol** *26*, 1728-1736.

van den Berghe V et al. (2013). Directed migration of cortical interneurons depends on the cell-autonomous action of Sip1. **Neuron** 77, 70-82.

School/Department:	Department of Cell biology, Erasmus MC
Supervisor information:	• Eskeatnaf Mulugeta, Ph.D., MSc., MBT., MBF., principal investigator,
	<u>e.mulugeta@erasmusmc.nl</u>
World no 30 Biomedical	• ORCiD: 0000-0003-4045-4835
<u>Sciences</u>	Website: <u>https://www.erasmusmc.nl/en/research/researchers/mulugeta-eskeatnaf</u>
	 Selected publication Blood, 2020 DOI: https://doi.org/10.1182/blood.2020004826 Cell Reports, 2020: DOI: https://doi.org/10.1016/j.celrep.2020.107647 Stem Cells, 2019: DOI: https://doi.org/10.1002/stem.3111 eLife, 2019 DOI: 10.7554/eLife.48561 Nature structural & molecular biology, 2019: DOI: https://doi.org/10.1038/s41594-019-0231-0 BioRxiv, 2017 DOI: https://doi.org/10.1101/209932 Genome research, 2016 DOI: https://doi.org/10.1038/nm.4098 Nature medicine, 2016 DOI: https://doi.org/10.1038/ncomms12222 Nature communications, 2016 DOI: https://doi.org/10.1038/ncomms12222 Nature, 2012: DOI: https://doi.org/10.1038/nature11070 Cell, 2009: DOI: https://doi.org/10.1016/j.cell.2009.10.034 Full list of publication: https://scholar.google.com/citations?hl=en&user=o5XA41sAAAAI
Project Title:	Systems Biology of Signaling and Transcription Factors
Abstract:	Cellular development and differentiation is a tightly controlled process that is orchestrated by the transcriptional regulation of genes. The control of gene transcription entails several layers of regulatory modules. Signaling pathways and their downstream TFs are important components of this gene transcription regulatory module and allow cells to properly respond to environmental cues. This interpretation within the cell's nucleus involves several genes that are organized in gene regulatory networks (GRNs), driving epigenomic and transcriptional changes and thereby cell fate, differentiation and maturation. We are interested in understanding the dynamics of such biochemical cascades and connected GRNs using in embryonic stem cells as a model. The aim of this PhD project is to understand the crosstalk and dynamics of signaling and TFs and their impact on the epigenome. To achieve this, we are using a holistic approach based on perturbation approaches and apply existing/emerging state-of-the-art computational and molecular biology techniques, including the development of novel single cell-omics techniques. Your responsibilities will include co-designing and performing such experiments, analyzing data, and documenting and reporting results in lab- and departmental meetings and at (inter-)national conferences
Requirements of candidate:	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills. Master degree or MD Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) English language requirement: English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs).

	Department of Cell Biology Erasmus MC	
Supervisor information:	Ana Ruiz-Saenz, Ph.D., principal investigator,	
	• Email: <u>a.ruizsaenz@erasmusmc.nl</u>	
World no 30 Biomedical	Website: https://www.erasmusmc.nl/en/research/researchers/ruiz-saenz	
Sciences	Grants:	
Sciences	H2020 Marie Skłodowska-Curie Individual Fellowship. (2020-2022)	
	AACR Scholar in Training Award (2017)	
	Post-doctoral Ramón Areces Foundation Grant (2013-2015)	
	EMBO Short-Term Fellowship (2009)	
	Most important publications:	
	- <u>Biochem Pharmacol.</u> (2021) doi: 10.1016/j.bcp.2020.114317.	
	 Mol Cancer Res (2021) doi: 10.1158/1541-7786.MCR-20-0825. Nature Cell Biology. (2019) doi: 10.1038/s41556-019-0328-z. 	
	- <u>Cell Reports</u> (2018) doi: 10.1016/j.celrep.2018.09.035.	
	- Cancer Research (2018) doi: 10.1158/0008-5472.CAN-18-0430.	
	- Journal of Clinical Oncology (2018) doi: 10.1200/JCO.2017.77.1899.	
	- <u>Breast Cancer Res Treat</u> . (2016) doi: 10.1007/s10549-016-3698-y.	
	 <u>Oncogene</u> (2015) doi: 10.1038/onc.2014.455. Journal of Cell Science (2013) doi: 10.1242/jcs.120840. Epub 2013 Aug 13. 	
	- Journal of Cell Biology. (2012) doi: 10.1242/JCS.120840. Epide 2013 Adg 13.	
Project Title:	Exploring novel mechanisms of cancer progression in breast cancer	
Abstract:	Breast cancer has the highest mortality of any cancer in women worldwide. Over the last few	
	years, increased understanding of tumor biology has led to the development of targeted	
	molecular therapies, increasing survival and improving the quality of life. However, despite	
	these advances, resistance to therapies and cancer progression remain a burden in the	
	successful treatment of cancer. The molecular mechanisms driving resistance and cancer	
	progression are complex and encompass not only the cancer cell but its interaction with the	
	surrounding microenvironment. Our previous studies concentrated on the oncogenic function	
	of HER2 in HER2-amplified breast cancers (<i>Cancer Research</i> 2018) and a new strategy to target	
	the undruggable HER3 (Oncogene 2015).	
	Recent studies of tumor genomes have identified mutations in novel genes without clear links	
	to cancer. We are particularly interested in deciphering the impact that those mutations have in	
	cancer progression and response to treatment. In this context, your work will focus on	
	unraveling novel mechanisms of genetic deregulation in cancer progression in collaboration	
	with other groups at the Medical Oncology and Cell Biology Departments. The work	
	encompasses a wide range of experimental techniques including protein biochemistry and cell	
	signaling, gene expression regulation, CRISPR technology, and interrogation of clinical samples.	
	Your responsibilities will include co-designing and carrying out experiments, analyzing data, and	
	documenting and reporting results in lab and departmental meetings.	
	We aim to create and foster a professional, creative, inclusive and productive environment,	
	where all lab members are empowered with the skills, knowledge and resources required for	
	their projects and future careers. To do so, team members are expected to be ambitious, critical	
	and take full responsibility for their projects in a supportive, collaborative and open culture.	
Requirements of	• We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in	
candidate:	using team work to tackle large scientific questions and thus requires a student with good communication skills.	
canalaate.	Master degree or MD	
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) 	
	English language requirement:	
	 English speaking countries & Netherlands: no requirement 	

School/Department:	Department of Cell Biology Erasmus MC	
Supervisor information:	• Jeffrey van Haren, Ph.D., principal investigator,	
	Email: <u>a.vanharen@erasmusmc.nl</u>	
World no 30 Biomedical	Website: <u>https://www.erasmusmc.nl/en/research/researchers/haren-jeffrey-van</u>	
<u>Sciences</u>	Grants/ awards:	
	- H2020 Marie Skłodowska-Curie IF(2020-2022)	
	 ASCB/EMBO Travel Award Postdoctoral Fellows (2017) Selected publications: 	
	 <u>Nature Cell Biology</u> (2018) doi: 10.1038/s41556-017-0028-5 <u>Current Opinion in Cell Biology</u> (2020) doi: 10.1016/j.ceb.2020.03.003 	
	- Journal of Cell Biology (2021) doi: 10.1083/jcb.201905199	
	- <u>Current Biology</u> (2016) doi: 10.1016/j.cub.2016.04.020	
	- <i>Current Biology</i> (2014) doi: 10.1016/j.cub.2014.06.037	
	- Genes and Development (2013) doi: 10.1101/gad.216200.113	
	- <u>Cell Reports</u> (2012) doi: 10.1016/j.celrep.2012.08.040	
	 <u>Molecular Biology of the Cell</u> (2010) doi: 10.1091/mbc.E09-12-1036 <u>Current Biology</u> (2010) doi: 10.1016/j.cub.2010.04.024 	
	- <u>Journal of Cell Biology</u> (2008) doi.org/10.1083/jcb.200707203	
Project Title:	Understanding directional neuronal migration in the developing nervous system	
-		
Abstract:	This project aims at improving our understanding of the cellular machinery that controls	
	neuronal migration and guidance. Defective guidance of neurons during embryonic	
	development leads to various neuro-developmental disorders such as lissencephaly, Joubert	
	syndrome, Hirschsprung's disease, and dysgenesis of the corpus callosum, and is linked to	
	autism and epilepsy. Long range guidance of neurons involves the detection of guidance	
	molecules, secreted by cells at a distance. Concentration gradients of such molecules can either	
	attract or repel neurons (a process termed chemotaxis). While many guidance signals and their	
	receptors have been identified, it is still largely unclear how guidance signals are processed in	
	space and time within the neuron, and how such signals direct localized assembly/disassembly	
	of the actin cytoskeleton, which is the main driver of cell movement. To improve our	
	understanding of this process, we will construct a parts list of the neuronal guidance machinery,	
	and observe/quantify the dynamics of these components using advanced microscopy	
	approaches. Furthermore, we will utilize novel approaches to control cell guidance, and aim to	
	develop a highly standardized <i>in vitro</i> neuronal guidance assay that will allows us to perform	
	quantitative analysis of cytoskeletal assembly in navigating cells. This project involves a wide	
	range of experimental techniques including novel optogenetic perturbation techniques (see	
	NCB 2018 , doi:10.1038/s41556-017-0028-5), live cell microscopy (including spinning disk	
	confocal microscopy, TIRFM and LLSM), CRISPR, micropatterning and protein engineering.	
	Knowledge gained from these studies will improve our understanding of human neuronal	
	migration / guidance disorders, and might in the future help in regenerative medicine, or the	
	development of advanced organ-on-chip technology. Your responsibilities will include co-	
	designing and performing experiments, analyzing data, reporting/presenting results (e.g. in lab	
	meetings, graduate school events, and at (inter-)national conferences), and in collaboration	
	with Dr. van Haren write research manuscripts.	
Requirements of	• We are looking for a highly motivated, hardworking student with a background in molecular cell biology,	
candidate:	nanobiology or related fields to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills.	
	 Successful applicants are expected to be ambitious, critical thinkers who take responsibility for their 	
	projects in a supportive, collaborative and open culture.	
	 Master degree or MD, preferably with experience in basic molecular biology techniques. 	
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could 	
	help with the scientific part of your scholarship proposal)	
	English language requirement:	
	• English speaking countries & Netherlands: no requirement	
	• Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)	

Department of Child & Adolescent Psychiatry

School/Department:	Department of Child and Adolescent Psychiatry, collaborating Department: Department of
	Epidemiology, Erasmus MC
Supervisor information:	Prof. dr. Henning Tiemeier
	Email: <u>h.tiemeier@erasmusmc.nl</u>
world no 28 in Social Sciences	Website: https://www.hsph.harvard.edu/henning-tiemeier/
& Public Health	Grants: multiple EU-Horizon2020 grants, NIH-NICHD grant, both VIDI and VICI, (see
	https://www.nwo.nl/en/researchprogrammes/nwo-talent-programme), EU Norface grant
world no 58 in	one of the world's 165 most highly cited scientists in the field of Social Science, general
Psychiatry/Psychology	(Clarivate/Thompson Reuters 2017, 2018 and again in 2019) H-index: 92 (Web of Science), 127 (Google Scholar)
<u> </u>	 Most important publications: KW Jansen TA, Korevaar TIM, Mulder TA, White T, Muetzel RL, Peeters RP, Tiemeier H. The Association of Maternal Thyroid Function during Pregnancy with Child Brain Morphology: A Time Window-Specific Analysis in a Prospective Cohort Study. Lancet E&D 2019; 7:629-637.
	 Xerxa Y, Delaney SW, Rescorla LA, Hillegers MHJ, White T, Verhulst FC, Muetzel RL, Tiemeier H. Association of Poor Family Functioning From Pregnancy Onward With Preadolescent Behavior and Subcortical Brain Development. JAMA Psychiatry. 2021;78(1):29-37.
	 Zou R, Tiemeier H, van der Ende J, Verhulst FC, Muetzel RL, White T, Hillegers M, El Marroun H. Exposure to Maternal Depressive Symptoms in Fetal Life or Childhood and Offspring Brain Development: A Population-Based Imaging Study. Am J Psychiatry. 2019; 176:702-710.
	 Rietveld CA, Medland SE, Derringer J, Yang J, Esko T, Martin NW, Westra HJ, Shakhbazov K, Abdellaoui A, () Teumer A; LifeLines Cohort Study, Tiemeier H, van Rooij FJ, Van Wagoner DR, Vartiainen E, Viikari J, Vollenweider P, Vonk JM, Waeber G, Weir DR, Wichmann HE, Widen E, Posthuma D, van Duijn CM, Visscher PM, Benjamin DJ, Cesarini D, Koellinger PD. GWAS of 126,559 individuals identifies genetic variants associated with educational attainment. Science. 2013;340:1467-71.
Project Title:	Early life adversity, maternal psychopathology, parenting and offspring
	neurodevelopment
Abstract:	Project Background: Many children experience early life adversities such as poverty,
	inadequate housing, poor neighbourhood, or parental psychopathology. These adversities
	have been repeatedly related to less optimal child development. What is less know are the
	protective factors that provide resilience against adversity, in particular whether supportive
	parenting, good family functioning or peer friendships provide buffering against the impact of
	adversity on behaviour and cognition. Also, in this project repeated brain imaging measures in
	adolescence will enable us to identify whether the interplay of childhood adversity and
	buffering factors impacts brain development in adolescence.
	<u>Aim:</u> The student will investigate how potential resilience or buffering factors, i.e. supportive
	parenting, neighborhood safety and peer friendship protect against poor behavioral and cognitive outcomes in children with and without experience of adversity.
	<u>Study Design and Methods:</u> The Generation R Study is a population-based cohort. Behavioral
	and brain imaging assessment at 10 and 13 years has been completed. Adversities such as
	parenting have been observed and assessed by questionnaire, father and mother mental
	health has been studied from pregnancy onwards. Importantly, this project will utilize
	observations in the home setting conducted in about 4000 children in the first few months of
	life, peer ratings and community data on neighborhood health. Child behavioral problems
	were repeatedly measured by parent, teacher and self-report. Brain function and morphology
	assessments are available in $N \approx 5500$ children and adolescents.
	Training in neuroscience and epidemiology leading to a MSc Epidemiology from Netherlands
_ _ _ _ _ _ _ _ _ _	 Institute of Health Sciences (<u>https://www.nihes.com/</u>) is part of the PhD program. We are looking for a highly motivated student to join our very international team. Our strength is in using team work to tackle
Requirements of candidate:	are scientific questions and thus requires a student with good communication skills.
	 Master degree or MD, background medicine, psychology, public health, epidemiology or neuroscience Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal)
	• English language requirement:
	• English speaking countries & Netherlands: no requirement
	O Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

Department of Clinical Genetics

The department Clinical Genetics performs innovative and high quality scientific research with a focus on three cornerstones: neurogenetics; genetics of congenital anomalies and genetics of cardiovascular disorders. The research focusses on both fundamental research to understand the mechanisms which cause hereditary diseases, as well as translational research for a quick translation of knowledge and renewing technology to improve diagnoses and treatments in favor of patients.

Some examples of diseases that are studied within our research section are: Fragile X syndrome, Parkinson disease, FXTAS, white matter disorders, malformations of cortical brain development, Hirschsprung disease and Pompe disease. Recently, three new research lines have been started focused on 1) aneurysms 2) the role of microglial cells in neurological diseases and 3) the role of the non-coding genome in gene regulation and genetic disorders. Additional research lines include: research om human cancers (uveal melanoma, Lynch Syndrome, breast cancer), psychological aspects of prenatal genetic testing and Non Invasive Prenatal Testing (NIPT).

We use state of the art methods to studying hereditary monogenic and polygenic disorders. Next Generation Sequencing and functional studies play an important role in unraveling disease mechanisms. For functional genetics and genomics, *in vitro* as well as *in vivo* models are used. We apply state-of-the-art methodologies, such as the use of induced pluripotent stem cells (so-called iPS-cells) generated from patients, disease modelling of brain development using cerebral organoids and epigenome characterization using massively-parallel-reporter assays. Widely applied animal models for the functional research are genetically modified mice and zebrafish. The functional work is performed in close cooperation with the Functional Unit of the Diagnostic section and the counseling section through which patients can be recruited. Currently, approximately 70 people are working in the research section, among which 30 PhD students. Most of these people are paid by external funding from many different funding bodies such as the EU, NIH, NWO, ZonMW, KWF, Heart foundation, Parkinson Foundation META kids and the Brain and Behaviour Research foundation.

On our website the different research lines are described in more detail

https://www.erasmusmc.nl/klinische_genetica/research/lijnen/

Our Principal Investigators (PIs) can be found on:

https://www.erasmusmc.nl/klinische_genetica/research/introduction/

A film presenting several of the research line can be found on:

https://www.youtube.com/watch?v=7iYn9DaCmbA&feature=youtu.be

Selection of recent publications

- Qaudri M et al. LRP10 genetic variants in familial Parkinson's disease and dementia with Lewy bodies: a genome-wide linkage and sequencing study. Lancet Neurol. 2018 17(7):597-608
- Tedja MS, et al. Genome-wide association meta-analysis highlights light-induced signaling as a driver for refractive error... NatureGenetics 2018;50(6): 834-848.
- Barakat TS, et al., Functional Dissection of the Enhancer Repertoire in Human Embryonic Stem Cells. Cell Stem Cell. 2018; Aug 2;23(2):276-288.e8.
- Oosterhof N, et al. Colony-Stimulating Factor 1 Receptor (CSF1R) Regulates Microglia Density and Distribution, but Not Microglia Differentiation In Vivo. Cell Rep 2018 24(5):1203-1217
- Bergsma AJ, et al., Alternative Splicing in Genetic Diseases: Improved Diagnosis and Novel Treatment Options. Int Rev Cell Mol Biol. 2018;335:85-141.
- van Poppelen NM, et al., Genetic Background of Iris Melanomas and Iris Melanocytic Tumors of Uncertain Malignant Potential. **Ophthalmology. 2018**, pii: S0161-6420(17)32844-0.
- van der Steen SL, et al., Choosing between Higher and Lower Resolution Microarrays: do Pregnant Women Have Sufficient Knowledge to Make Informed Choices Consistent with their Attitude? J Genet Couns. 2018;27(1):85-94.
- van Waning JI, et al. Genetics, Clinical Features, and Long-Term Outcome of Noncompaction Cardiomyopathy. J Am Coll Cardiol. 2018, 71(7):711-722
- Halim D, et al. Loss of LMOD1 impairs smooth muscle cytocontractility and causes megacystis microcolon intestinal hypoperistalsis syndrome in humans and mice. **Proc Natl Acad Sci U S A. 2017**, 114(13):E273.
- Olgiati S, et al., DNAJC6 Mutations Associated With Early-Onset Parkinson's Disease. Ann Neurol. 2016; 79(2):244-56.
- Zeidler S, et al., Combination Therapy in Fragile X Syndrome; Possibilities and Pitfalls Illustrated by Targeting the mGluR5 and GABA Pathway Simultaneously. Front Mol Neurosci. 2017;10:368.
- Goverde A et al., Small-bowel Surveillance in Patients With Peutz-Jeghers Syndrome: Comparing Magnetic Resonance Enteroclysis and Double Balloon Enteroscopy. J Clin Gastroenterol. 2017;51(4):e27-e33.

Department of Clinical Genetics

School/Department:	Department of Clinical Genetics Erasmus MC	
Supervisor information:	Stefan Barakat, M.D., Ph.D., MSc., principal investigator	
	Email: <u>t.barakat@erasmusmc.nl</u>	
World no 30 Biomedical	Website: <u>https://www.erasmusmc.nl/en/research/groups/barakat-lab-non-coding-genome-in-clinical-genetics</u>	
<u>Sciences</u>	Personal Grants: Nick Starson Followship (2014): EMBO Long Term Followship (2014): Mario Składowska Curio Individual Followships (JE EE)	
	 Niels Stensen Fellowship (2014); EMBO Long-Term Fellowship (2014); Marie Skłodowska-Curie Individual Fellowships (IF-EF) (2015); Human Frontiers Science Project Long-Term Fellowship (2015); Wellcome Trust ISSF2 award (2015); NARSAD Young Investigator Award (2016); ZonMW VENI award (2016); Erasmus MC fellowship (2017); EMC Human Disease Model Award (2018) Awards: 	
	American Society of Human Genetics (ASHG) Charles J. Epstein Award for Excellence in Human Genetics Research (2015); International Society for Differentiation Beverly Kerr McKinnel Award, for outstanding research as a PhD student (2012)	
	Most important publications: (H-index:14; total citations:>1320)	
	(sep 2020) Nature Reviews Neurology doi: 10.1038/s41582-020-0395-6 (IF: 27.0) (apr 2020) Acta Neuropathologica doi: 10.1007/s00401-020-02128-8 (IF18.2)	
	(der 2019) Acta Neuropathologica doi: 10.1007/s00401-020-02128-8 (IF18.2)	
	(aug 2018) Cell Stem Cell doi: 10.1016/j.stem.2018.06.014 (IF:23.3)	
	(aug 2015) Genome Biology doi: 10.1186/s13059-015-0698-x (IF:11.9)	
	(mar 2014) Molecular Cell doi: 10.1016/j.molcel.2014.02.006 (IF:14.7) (mar 2013) Cell Reports doi: 10.1016/j.celrep.2013.02.018 (IF:8.3)	
	(apr 2012) Nature doi: 10.1038/nature11070 (IF:40.1)	
	(jun 2012) Molecular Cell doi: 10.1016/j.molcel.2012.04.003 (IF:14.7)	
	(oct 2011) Nucleic Acid Research doi: 10.1093/nar/gkr550 (IF:9.2) (jun 2010) Cell Stem Cell doi: 10.1016/j.stem.2010.05.003 (IF:23.3)	
	(nov 2009) Cell doi: 10.1016/j.cell.2009.10.034 (IF:30.4)	
	For full list see: <u>https://www.ncbi.nlm.nih.qov/pubmed/?term=tahsin+stefan+barakat</u>	
Project Title:	Deciphering the role of Non-Coding DNA sequences in the genetics of	
	neurodevelopmental disorders	
Abstract:	Despite the fact that we know that the majority of DNA sequences (~98%) in the human genome do	
	not encode protein-coding genes, our understanding of those sequences and why they are important	
	is still far from complete. An important group of non-coding genome elements are enhancers that	
	are crucial for the proper regulation of spatiotemporal gene expression. The clinical genetic work-up	
	of patients suffering from neurodevelopmental disorders currently focusses almost completely on	
	exons. An attractive hypothesis is that currently genetically unexplained patients might have	
	mutations in regulatory elements such as enhancers that might cause their phenotype, but before	
	this hypothesis can be tested on a large scale it is crucial to identify regulatory elements involved in brain development.	
	In my lab, we are trying to understand the role of regulatory elements in brain development using	
	several approaches. We are using state-of-the-art techniques to profile the epigenome of cerebral	
	organoids using ChIP-seq, ATAC-seq, and single cell RNA-seq to identify putative regulatory	
	elements. Using ChIP-STARR-seq, a novel type of massively parallel reporter assay system that we	
	have developed, we are generating genome-wide enhancer activity maps of various brain related	
	cell types. Using functional genomics and CRISPR-Cas9 mediated screens, we validate putative	
	enhancers. Integrative computational analysis and data mining further helps us to identify crucial	
	regulatory elements, that we sequence in a large cohort of genetically unexplained patients. Using	
	iPSC technology combined with genome-engineering, we validate our findings. In addition, we	
	perform disease modeling for novel genetic neurodevelopmental disorder. Ultimately, our efforts	
	will lead to an enhanced understanding of the brain regulome and will lead to novel diagnostic	
	approaches for patients suffering from neurodevelopmental disorders.	
Requirements of	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills. 	
candidate:	 Master degree or MD 	
	Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the	
	scientific part of your scholarship proposal)	
	English language requirement:	
	English speaking countries & Netherlands: no requirement	

Department Clinical Genetics

School/Department:	Department of Clinical Genetics, Erasmus MC	
Supervisor information: World no 30 Biomedical Sciences	 Prof. dr. Ype Elgersma, <u>v.elgersma@erasmusmc.nl</u> Websites: <u>www.neuro.nl/research/elgersma</u> <u>www.encore-expertisecentrum.nl</u> <u>www.functionalgenomics.nl</u> Personal Grants: VIDI, VICI Most important publications: Mol Psych 2015 20:1311-21 JAMA Neurology 2015: 72:1052–1060. Nature 2015 526:50-1 J Clin Invest 2015 125:2069-2076 Am J Hum Genet 2017 5:768-788 Mol Psych 2019 24: 757-771 Nature Neuroscience 2019 22:1235-1247 	
Project Title:	Gaining insight in the molecular mechanisms underlying neurodevelopmental disorders.	
Abstract:	 Neurodevelopmental disorders (i.e. intellectual disability, autism) affect >1% of the population, and often have a genetic basis. Our lab seeks to get insight in the molecular and cellular mechanisms underlying these disorders, with the ultimate goal to develop treatments. Our research into these disorders is divided into three research lines: (1) Improving genetic diagnosis, (2) Understanding the mechanisms underlying neurodevelopmental disorders, and identifying treatments (3) Translational studies (i.e. clinical trials) to improve the quality of life of the affected individuals. For the candidate student we have possibilities to join the following projects: Improving diagnosis: To improve genetic diagnosis, we have developed a functional genomics screen (PRiSM) (see functionalgenomics.nl) to rapidly determine if a genetic variant is pathogenic. This screen is not only important for providing a diagnosis, but also allows us to get more insight in the genes underlying neurodevelopment. New assays will be developed and validated for this screen. Understanding the mechanisms and identify treatments: To get more insight in the pathophysiology of neurodevelopmental disorders, we typically make use of genetically engineered mouse models as a tool to dissect the underlying mechanisms. Mouse models are analyzed at the biochemical, cellular (electrophysiological) and behavioral level. By analyzing the mice at all these levels we hope to understand the specific function of these genes and proteins in brain development and learning and memory. Besides mouse models, we are also using iPS cells to study these disorders. The genes and proteins that we in particular focus on are proteins associated with the RAS-ERK-MTOR signaling pathway and the proteasome. Treatments that we are in particular interested in are antisense oligonucleotide (ASO) treatments, that target directly the mutated RNA. 	
Requirements of candidate:	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills. Master degree or MD Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) English language requirement: English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs) 	

School/Department:	Department of Epidemiology Erasmus MC	
Supervisor information:	Dr. Maryam Kavousi, Associate Professor	
	Email: <u>m.kavousi@erasmusmc.nl</u>	
World no 21 Public,	Website: <u>http://www.erasmus-epidemiology.nl/</u>	
Environmental &	Personal Grants: AVA Descent 2012	
Occupational Health	 AXA Research Grant, 2012 IDF, 2014 	
	Prestigious UNESCO-Loreal Fellowship 'For Women in Science', 2014	
	Prestigious ZonMw VENI Grant, 2015	
	 Erasmus MC Mrace Grant, 2016 ZonMw Grant, 2017 	
	Hartsticthing (Dutch Heart Foundation) Grant, 2017	
	Most important publications:	
	 Nature Genetics 2011 43(10):940-947 	
	 Circulation 2011 124(25):2855-2864 	
	• Circulation 2012 126(4):468-478	
	• Annals of Internal Medicine 2012 156(6):438-444	
	• JAMA 2014 311(14):1416-1423	
	• JAMA 2016 315(23):2554-2563	
	 JAMA Cardiology 2016 1(6):708-713 	
	 JAMA Cardiology 2016 1(7):767-776 	
	• JAMA 2016 316(20):2126-2134	
	 JAMA Cardiology 2017 2(9):986-994 	
	• Circulation Research 2017 121(12):1392-1400	
	• Nature Genetics 2018 50(9):1225-1233	
Project Title:	Global Cardiomtabolic Risk Profile	
Abstract:	Population aging is magnifying the global burden of cardiometabolic disorders and their consequences.	
	Global cardiometabolic risk represents the overall risk of developing cardiovascular diseases and/or type	
	2 diabetes due to a cluster of risk factors. Development of clinically useful primary and secondary	
	prevention strategies will require a more comprehensive understanding of these complex conditions. We	
	study the association of traditional and novel risk factors, representing of different pathophysiologic	
	pathways, with cardiometabolic risk across its spectrum. The risk factors comprise biomarkers, including	
	the novel omics markers, as well as the new cardiovascular imaging markers.	
	Besides contribution of various pathways, as well as their interactions, to form the natural course of cardiometabolic disorders, differences between women and men in these processes are highly of interest.	
	The studies are performed within the Cardiometabolic research line of the Department of Epidemiology	
	using the large population-based Rotterdam Study. We closely collaborate with other renowned	
	population-based studies across Europe and United States including the cohorts involved in the	
	international CHARGE Consortium (The Cohorts for Heart and Aging Research in Genomic Epidemiology).	
Requirements of	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills. 	
candidate:	 Master degree or MD 	
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific next of your scholarship menoce) 	
	 scientific part of your scholarship proposal) English language requirement: 	
	English speaking countries & Netherlands: no requirement	
	Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)	

Supervisor Information: Dr. Daniel Bos, MD, PhD E- Indit: d. bost@crosumes.nl Headit: d. bost@crosumes.nl	School/Department:	Department of Epidemiology, Erasmus MC	
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Requirements of • We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using			
toom work to tackle large ccientific substitute and thus requires a student with good communication shills	Requirements of		
candidate:team work to tackle large scientific questions and thus requires a student with good communication skills.Master degree or MD	candidate:		thus requires a student with good communication skills.
 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the 		u	lowance and international air plane ticket (we could help with the
scientific part of your scholarship proposal)			
English language requirement:			
 English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs) 			

School/Department:	Department of Epidemiology, Erasmus MC	
Supervisor information:	Dr. Mohsen Ghanbari	
	Assistant professor, Principal investigator of the Molecular & Systems Epidemiology group	
<u>World no 21 Public,</u>	Email: m.ghanbari@erasmusmc.nl	
Environmental &	Website: <u>http://www.erasmus-epidemiology.nl</u>	
Occupational Health	https://www.erasmusmc.nl/en/research/researchers/ghanbari-mohsen	
	• Grants:	
	 Early Career Award, The Cohorts for Heart and Aging Research in Genomic Epidemiology, 2018 European Foundation for the Study of Diabetes Fellowship, 2018 	
	 Alzheimer Netherland Fellowship, 2018 	
	Most important publications:	
	Dr. Ghanbari has so far published over 80 international peer-reviewed publications.	
	Nature Communications. 2021 May 14;12(1):2830. Epigenome-wide association meta-analysis of	
	 Stroke. 2021 Mar;52(3):945-953. Circulatory MicroRNAs as Potential Biomarkers for Stroke Risk Brain. 2020 Apr 1;143(4):1220-1232. Plasma tau, neurofilament light chain and amyloid-β levels 	
	 Cell. 2020 Sep 3;182(5):1214-1231. The Polygenic and Monogenic Basis of Blood Traits and Diseases. 	
	 Diabetes Care. 2020 Apr;43(4):875-884. Epigenetic Link Between Statin Therapy and Type 2 Diabetes. 	
	• Nature Communications. 2019 Aug 20;10(1):3346. A metabolic profile of all-cause mortality risk	
	• Human Mutation. 2019 Nov;40(11):2131-2145. A functional variant in the miR-142 promoter	
	• Nature Genetics. 2019 Apr;51(4):636-648. Multi-ancestry genome-wide gene-smoking interaction	
	Nature Communications. 2019 Jan 22;10(1):376. Multi-ancestry study of blood lipid levels identifies	
	• Gastroenterology. 2017 Oct;153(4):1096-1106. Epigenome-Wide Association Study Identifies	
Project Title:	Integration of population-based omics data to explore molecular mechanisms	
	underlying age-related diseases	
Ale adversate		
Abstract:	Genetic and molecular epidemiology are emerging innovative fields of research in which	
	molecular and biological concepts are incorporated into computational models and	
	epidemiologic studies to identify genetic predispositions of complex diseases. This is made	
	possible by recent rapid technological advances in high-throughput laboratory assays that	
	measure various biomarkers from biological samples. Although traditional epidemiology has	
	been proven valuable to identify associations between exposure and disease in populations;	
	yet, it does so without obtaining information of the biological processes that underlie the	
	associations. Molecular epidemiology could enhance the measurement of exposure, effect,	
	and susceptibility, and give insight into biological mechanisms. This knowledge will ultimately	
	lead to the identification of early etiologic, diagnostic, and prognostic markers of diseases,	
	allow us to better target preventive strategies and yield new therapeutics for complex	
	diseases.	
	Within the Molecular & Systems epidemiology research line of the department of	
	Epidemiology, we conduct cutting-edge research on the genetic determinants and novel	
	biomarkers of age-related diseases (e.g., Cardiovascular disease, type 2 diabetes, Alzheimer's	
	disease, fatty liver disease) using omics data (incl. genomics, epi-genomics, transcriptomics,	
	proteomics, and metabolomics) from the Rotterdam Study, a large population-based cohort	
	of 15,000 participants followed since 1990. Moreover, we closely collaborate with several	
	renowned international population-based cohort studies across Europe and United States on	
	large-scale international projects.	
Requirements of	• We are looking for a highly motivated, bright student to join our international and multidisciplinary team. For	
candidate:	this projects, using big data and often collaborating in consortia, we require strong statistical skills and good	
	communication skills.	
	• The student should have an MD or Master degree in Biology, Epidemiology, Biostatistics or a related field, and	
	should be fluent in English (IELTS≥7.0 (≥6.0 for all subs), TOEFL ≥100 (≥20 for all subs).	
	We offer: Supervision, data access, advanced courses in genetic epidemiology and biostatistics, research	
	infrastructure, and other training. Your salary and living expenses should be covered by the scholarship. We	
	could help with the scientific part of the proposal. For more information related to this proposal, please contact	

School/Department:	Department of Epidemiology, Erasmus MC	
Supervisor information: World no 21 Public, Environmental & Occupational Health	 Prof dr M. Kamran IKRAM Email: m.ikram@erasmusmc.nl Website: https://www.erasmusmc.nl/en/research/departments/epidemiology Grants: Lee Kuan Yew Fellowship, Singapore (2011) VENI, Netherlands Organisation for Scientific Research, the Netherlands (2012) National University Health System, National University of Singapore, Clinician Scientist Program Grant, Singapore (2012) National Medical Research Council, Clinician Scientist Award, Investigator Category, Singapore (2013) European Institute of Innovation and Technology (2016) ParkinsonFonds, the Netherlands (2018) Netherlands Organization for Scientific Research – Covid 19 Program, the Netherlands (2020) Most important publications: Mov Disord 2020; Sept 23 Epub J Am Coll Cardiol 2020; 75:2387-2399 Plas Med 2019;16:e1002933 Nat Genet 2019;51:1624-1636 Nature Medicine 2019;25:1364-1369 Circulation 2019;139:1698-1709 Int J Epidemiol 2019;48:1286-1293 JAMA Neurol 2018;17:434-444 Circulation 2017;135:2207-2209 	
Project Title:	Nat Neurosci 2016;19:1569-1582 Nature 2016;536:41-47 Vascular disease and autonomous dysregulation in Parkinson's Disease	
Abstract:	Parkinson's disease (PD), which is the most common subtype of parkinsonism, is a chronic neurodegenerative condition in the elderly. Although several environmental and genetic factors have been implicated in the development of parkinsonism, there is still uncertainty about the exact mechanisms underlying neuronal cell loss in these conditions. Among others, a potential role of vascular disease has been hypothesized based on the observation that that markers of vascular pathology are strongly related to two other common neurological syndromes, namely stroke and dementia. Furthermore, a high prevalence of lacunar infarcts in the basal ganglia of patients with parkinsonism have been reported. During the course of dementia 25% of patients develop parkinsonism, whereas approximately a third of patients with PD are eventually diagnosed with dementia. However, in spite of an overlap in clinical and pathological features between these neurological syndromes, the role of vascular pathology in the etiology of parkinsonism syndromes remains unclear. Besides vascular disease, cardiovascular dysregulation, as a manifestation of autonomous dysfunction, has also been implicated in PD. However, these observations have mainly come from clinical studies, in which the exact order of events is difficult to disentangle (reverse causality). Thus far, observations from population-based studies are largely lacking. In view of these gaps in the literature, our overall aim of this project is to determine the role of vascular disease and autonomous dysfunction in the development of Parkinson's disease and non- PD parkinsonism. To accomplish this data from the large population-based Rotterdam Study (N=14,926), which has been running for more than 30 years, will be used. Within this cohort, extensive cardiovascular risk factors assessment, including imaging of the major arteries in the heart-brain axis, has been performed. All persons are also evaluated for parkinsonism, using questionnaires, extensive examinations at our research center a	
Requirements of candidate:	 team. Due to the nature of the project and data, strong statistical skills and good communication skills are required. The student should have completed an MD or MSc in Neurosciences, Psychology, Health Sciences, Epidemiology, or a related field. A good command of English is required (level of IELTS 7.0 (min 6.0 for all subs) or TOEFL 100 (min 20 for all subs). Within the project the student will have access to the Rotterdam Study data, training in epidemiology and statistics, and the broader Erasmus MC research infrastructure. The scholarship will, at least, have to cover subsistence allowance and international air plane ticket. We are happy to help with the scientific part of your scholarship proposal, please contact prof.dr. M.K. Ikram (m.ikram@erasmusmc.nl) 	

School/Department:	Department of Epidemiology Erasmus MC	
Supervisor information:	Prof.dr. M. Arfan Ikram; dr. Gennady Roshchupkin	
	• Secondary affiliation: Adj. professor at Harvard Chan School of Public Health, Boston (MAI)	
World no 21 Public,	Email: <u>m.a.ikram@erasmusmc.nl</u> ; <u>g.roshchupkin@erasmusmc.nl</u>	
Environmental &	Website: <u>https://www.erasmusmc.nl/en/research/researchers/ikram-arfan-m</u>	
Occupational Health	Personal Grants: The leasanth for diagonal left 10 years is more than 15 ME we including EBC Starting Creat European IBND screet	
	Total research funding over last 10 years is more than 15 MEuro, including ERC Starting Grant, European JPND grant, multiple Horizon 2020 consortium collaborations, multiple NIH R01-subcontract PI.	
	He has supervised 28 PhD students.	
	Most important publications:	
	Satizabal CL. Nat Genetics 2019	
	Wang J. PNAS 2019 Hibar DP. Nat Commun 2017	
	Adams HH. Nat Neurosc 2016	
	Roshchupkin GV. Nat Commun 2016	
	Ikram MA. Nat Genetics 2012 Ikram MA. NEJM 2009	
Project Title:	Deep Learning in Omics Data Analysis and Precision Medicine	
Abstract	A central goal of human genetics is to understand the relationship between genetic variation	
	and diseases or traits. There are many different technologies, study designs and analytical tools	
	for identifying such relations. Recent technological advances and biobank initiatives have	
	allowed studies involving hundreds of thousands, and even millions, of individuals. Moreover,	
	many studies have started collected other omics data beyond genetic data, including gene	
	expression, methylation, proteins, metabolites, and microbiome. This allows getting closer to	
	the trait's etiology. However, by nature most of the analytical tools and methods are either	
	univariate or cannot handle multi-omics data. Therefore, cross-omics methods are missing.	
	Human genetics needs new types of approaches to solve such problems for improving the	
	diagnosis, treatment, and classification of complex diseases.	
	Deep learning (DL) is a rapidly growing field. The application of the neural networks has become	
	a golden standard in many research areas. DL algorithms have shown successful ability to detect	
	a complex pattern in high-dimensional data, and also are able to integrate data from various	
	resources by having many input channels into neural network	
	The main goal of this project is to develop new DL methods for multi-omics analysis, which	
	will be able to integrate prior biological knowledge and improve our understanding of the	
	etiology of complex traits, such as dementia and cognition. An additional dimension in this	
	project will be to combine the various omics data to brain MRI-imaging. We aim to apply these	
	methods on large datasets from population-based Rotterdam study, UK Biobank as well as	
	within international CHARGE consortium.	
Requirements of	We are looking for a highly motivated, hardworking student to join our very international team. Successful candidates are expected	
candidate:	to have a strong quantitative or computer science background, excel at critical thinking, with a strong motivation to engage in the development and application of advanced analytical methods. The following are strongly preferred requirements for interest	
	candidates:	
	• Master degree in mathematics, computer science, statistics, bioinformatics, physics, electrical engineering, or in an equivalent	
	 discipline. Strong knowledge of Python and R. 	
	•Experience with machine learning and deep learning methods.	
	•Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part	
	of your scholarship proposal) •English language requirement:	
	English speaking countries & Netherlands: no requirement	
	- Other countries: IELTS 6.	
	We offer you: - Access to the research infrastructure at Frasmus MC (including Rotterdam Study and related datasets) as well as access	
	 Access to the research infrastructure at Erasmus MC (including Rotterdam Study and related datasets) as well as access to our network of international collaborations (>25 countries) 	
	- A dedicated team of supervisors (prof. Ikram dr. Roshchupkin) with longstanding expertise in epidemiology, -omics,	
	- A dedicated team of supervisors (prof. Ikram dr. Roshchupkin) with longstanding expertise in epidemiology, -omics, imaging, and deep learning	
	- A dedicated team of supervisors (prof. Ikram dr. Roshchupkin) with longstanding expertise in epidemiology, -omics,	

School/Department:	Department of Epidemiology, Erasmus MC	
Supervisor	Dr. Annemarie I. Luik, PhD	• Dr. Daniel Bos, MD, PhD
information:	Email: <u>a.luik@erasmusmc.nl</u>	Email: <u>d.bos@erasmusmc.nl</u>
	Website:	Website:
World no 21 Public,	https://www.erasmusmc.nl/en/research/groups/psychiatr	https://www.erasmusmc.nl/en/research/groups/imaging-
Environmental &	<u>ic-epidemiology</u>	<u>of-arteriosclerosis</u>
Occupational Health	Grants and Awards:	Grants and Awards:
	European Sleep Research Society Top Young Researcher	Royal Academy of Arts and Sciences Grant (2016)
	 Abstract (2018) Sleep Research Society Foundation Career Development 	Lourens Penning Prize for best publication in the field of Neuroradiology(2016)
	Award (2019)	 BrightFocus Foundation Grant (2017)
	• Netherlands Organization for Scientific Research (2020)	Erasmus MC Mrace Grant (2019)
	Most important publications:	• European Commission Horizon 2020 - Research and
	Nature Hum Behav 2020; in press.	Innovation Framework Programme (2019)
	• Mov Disord. 2020; published online Sep 15.	Netherlands Organisation for Scientific Research (2019)
	• Alzheimers Dement 2020; 16: 1259-1267.	Most important publications:
	• JAMA Psychiatry 2019; 76: 21-30.	• JACC 2020; 19;75:2387-2399.
	 JAMA Pediatrics 2019; 173: 883-885. Nature Genet 2019; 51: 387-393. 	 BMC Medicine 2020; 18:263. Heart 2020; 106(2):133-139.
	 Nature Comm 2019; 15: 1521. 	 Plos Med 2020; 17(5):e1003115.
	• Brain 2019; 142; 2013-2022.	 Eur Heart J 2018; 39:3369-3376.
	• NPJ Digital Med 2018; 1:3	• JACC 2018; 72: 582-584.
	• Lancet Psychiatry 2017; 4: 749-758.	• Alzheimers Dement 2018; pii: S1552-5260(18)30129-8.
	• Nature Genet 2017;49: 274-281.	• Eur Radiol 2018; 2018: 28:3082-3087.
	Psychol Med 2016; 46: 1951-1960.	• Circulation 2017; 135:2207-09.
	• Mol Psychiatry 2015; 20: 1232-1239.	• Circ Cardiovasc Genet 2013; 2013; 6:47-53.
Project Title:	Unravelling the role of vascular disease in d	epression
Abstract:	Depression remains one of the top causes of disabi	lity worldwide according to the World Health
	Organization. Interestingly, an increasing body of e	
		ent increase in the occurrence of depression around
	the age of 60 may even be largely attributed to vas	
		· ·
	relationship between vascular disease and depress	
		f research on vascular disease pertains to its location
	in the blood vessel system. Although vascular disea	
		rably across different blood vessels within the same
	person. As such, vascular disease located in the ma	in blood vessels that provide the brain with blood
	may thus play a more important role in the develop	amont of doprossion and doprossive symptoms than
	They thus pluy a more important role in the develop	sinent of depression and depressive symptoms than
	vascular disease in more distant arteries.	
	vascular disease in more distant arteries.	
	vascular disease in more distant arteries. The overall aim of this project is to comprehensivel	ly investigate the role of vascular disease in the
	vascular disease in more distant arteries. The overall aim of this project is to comprehensivel development of depression and to better understa	ly investigate the role of vascular disease in the nd the potential causal link between vascular disease
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Requirements of	 vascular disease in more distant arteries. The overall aim of this project is to comprehensivel development of depression and to better understa and depression. To accomplish this data from the la which has been running for more than 30 years, wi major arteries in the heart-brain axis has been perf depression, using questionnaires, clinical interview link between vascular disease and the developmen The studies will be performed within the Psychiatri and the Imaging of Arteriosclerosis research group Moreover, we participate in different large consort We are looking for a highly motivated, hardworking stuto the nature of the project and data, strong statistical imaging and mental health are required. The student should have completed an MD or MSc in Noreover is the student should have completed an MD or MSc in Noreover is the student should have completed an MD or MSc in Noreover is the project and data. 	ly investigate the role of vascular disease in the nd the potential causal link between vascular disease arge population-based Rotterdam Study (N=14,926), Il be used. Within this cohort, medical imaging of the formed. All persons are also extensively evaluated for s and follow-up of medical records. Henceforth, the t of depression can be established. c research group of the Department of Epidemiology of the Department of Epidemiology and Radiology. ita, including CHARGE and ENIGMA. udent to join our international and multidisciplinary team. Due skills, good communication skills, and an interest in medical Neurosciences, Psychology, Health Sciences, Epidemiology, or
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School/Department:	Department of Epidemiology, Erasmus MC
Supervisor information: World no 21 Public, Environmental & Occupational Health	 Dr.ir. Trudy Voortman Principal investigator Nutrition & Lifestyle Epidemiology, Life-course epidemiology Email: trudy.voortman@erasmusmc.nl Website: www.erasmusmc.nl/en/research/groups/nutrition-and-lifestyle-epidemiology; www.trudyvoortman.com Personal honors and grants: European Society for Clinical Nutrition and Metabolism (ESPEN) Fellowship 2020 American Society for Nutrition – Peter Reed Award for outstanding research in macronutrient metabolism, 2018 Thrasher Pediatric Medical Research Career Award, USA, 2016 European Foundation for the Study of Diabetes Fellowship, 2015 Selected member of the European Nutrition Leadership Platform (ENLP), 2015-present Most important publications: Dr. Voortman has published over 100 international publications, of which more than 60 publications as direct supervisor of the researchers in her team. Most PhD students in our team write 5 to 8 publications as first author within their PhD project and contribute to additional papers as coauthor. All publications in our team have been published in journals in the top quartile of their field and more than haf have been published in top-10% journals. BMJ-British Medical Journal 2017;356:j1000. Dairy consumption and risk of hypertension. Lancet 2018;391(10129):1513-23. Risk thresholds for alcohol consumption. The Lancet Diabetes & Endocrinology 2017;5(5):367-76. Vitamin D in pregnancy and child bone health Gastroenterology 2018; doi:10.1038/s41591-019-0547-7. Lifestyle and dementia risk. BMJ, 2019. doi: 10.1038/s41591-019-0547-7. Lifestyle and dementia risk. BMJ, 2019. doi: 10.1038/s41586-020-2338-1. Global repositionining of non-optimal cholesterol. Clinical
Project Title:	Nutrition and Lifestyle and cardiometabolic health across the life course: a focus on underlying pathways and mechanisms
Abstract:	Nutrition and lifestyle affect health throughout the life course: from pregnancy and infancy to old age. In our research group, we study nutrition and other lifestyle factors in pregnant women, children, adults and elderly; and how diet and lifestyle impact health in these groups. In these projects, we also focus on underlying mechanisms of how nutrition affects disease risk, including e.g. inflammation, metabolomics, DNA methylation, and gut microbiome composition. The studies are performed within the Nutrition & Lifestyle research group at the Department of Epidemiology, one of the world leading academic centers in epidemiology. The candidate can use data from large cohort studies available at the department and through collaborations in consortia. Studies at the department for example include the Rotterdam Study, a population based study among 15,000 people followed since 1990 and the Generation R Study, a birth cohort
	study in 10,000 mothers and their children. Our Nutrition & Lifestyle team closely collaborates with other research lines at Erasmus MC and other institutes across Europe and the United States, including the departments of Nutrition at Harvard School of Public Health, Wageningen University, Cambridge University, Tufts University.
	For more information about our team and department, please check our webpages: <u>www.erasmusmc.nl/en/research/groups/nutrition-and-lifestyle-epidemiology</u> and <u>https://www.erasmusmc.nl/en/research/departments/epidemiology</u>
Requirements of candidate:	 We are looking for a highly motivated student to join our very international and multidisciplinary team. For these projects, using large datasets and in collaborations with various other research groups, strong statistical and good communication skills are required. The candidate should have an MD or MSc degree in Health Sciences, Epidemiology, Biostatistics,. Nutrition Science, or a related field, and should be fluent in English (IELTS≥7.0 (≥ 6.0 for all subs), TOEFL ≥100 (≥ 20 for all subs).
	• We offer: Supervision by at least two supervisors, data access to cohort studies, advanced courses in epidemiology at our postgraduate research school NIHES, and other training. Your salary and living expenses should be covered by the scholarship. We are happy to discuss the details further with you directly and help with the scientific part of your proposal. Please contact dr. Trudy Voortman at trudy.voortman@erasmusc.nl

In a nutshell:

- Head: Prof. dr Marco Bruno
- Staff: 6 hepatologists, 10 gastroenterologists
- Trainees/fellows: 19 trainees, 2 foreign fellows for advanced training (6 months)
- GI translational lab: head Prof. dr Maikel Peppelenbosch
- 55 PhD students on liver, GI, clinical and/or translational projects
- GI clinical research unit: datamanagers, research nurses, statistician
- Current world ranking: no 14 (<u>US News subject ranking 2021</u>)

Well established interdisciplinary working relationships with department of surgery, oncology and radiology with both clinical and research activities being initiated and steered by multidisciplinary interest groups (liver centre, pancreas centre, esophageal cancer center

Clinical and translational research is centered around the following main topics:

Gastroenterology:

Oncology

o Pancreatic cancer (early diagnosis in high risk individuals, pancreatic cyst differentiation and follow-up, optimal palliative treatment strategies, neoadjuvant treatment in stage II/borderline disease, folfirinox followed by radiotherapy in locally advanced disease, pancreatic biopsies and personalized medicine)

o Esophagal cancer (neoadjuvant treatment strategies, Barrett's esophagus identification biomarkers for better risk profiling, drug prevention of Barrett's)

o Colonic cancer (colonic cancer in high risk populations, general population screening for colonic cancer) Advanced endoscopy

- o Resection techniques (EMR/ESD)
- o EUS (follow-up studies high risk pancreatic cancer, pancreatic cyst follow-up study, improving the yield of EUS-guided tissue sampling)
- o ERCP (stenting of benign biliary strictures with metal stents, biodegradable stenting of pancreatic strictures, advanced endoscopic imaging of biliary tree and pancreas, tissue sampling)
- o Esophagal stenting (optimal stent design and protocol in both malignant and benign strictures) *Inflammatory bowel disease*
- o Optimal en cost effective treatment with biologicals
- o IDB and pregnancy

Hepatology:

- o Viral hepatitis (novel treatment therapies, advanced imaging of the liver)
- o Cirrhosis (early detection of HCC, treatment of complications of portal hypertension)
- o Hepatocellular carcinoma (novel treatment strategies)risk profiling, prediction of response etc.

Publications, Grants:

See vacancy from the relevant PI

School/Department:	Department of Gastroenterology and Hepatology, Erasmus MC	
Supervisor information:	Andre Boonstra, PhD, Associate Professor - Immunology of Viral Hepatitis and Liver Cancer	
	Email: p.a.boonstra@erasmusmc.nl	
world no 14	For information about our research and laboratory: <u>www.viralhepatitis.nl</u> and https://www.erasmusmc.nl/en/research/groups/chronic-viral-hepatitis-liver-cancer	
Gastroenterology &	For information on our EU funded ESCALON project: <u>www.escalon.eu</u>	
<u>Hepatology</u>	Most relevant recent publications:	
	Hepatitis B core-specific memory B cell responses associate with clinical parameters in patients with chronic HBV. <u>J Hepatol. 2020</u> Jul;73(1):52-61.	
	Serum immune signatures associated with HCC development in DAA-treated HCV patients. <u>Gastroenterology. 2018. Feb;</u>	
	<u>154(3):515-517</u> .	
	Serum Biomarkers for the Prediction of Hepatocellular Carcinoma. <u>Cancers. 2021; 13(7):1681</u> Hepatitis B core-related antigen levels predict recurrence-free survival in patients with HBV-associated early-stage hepatocellular	
	carcinoma: results from a Dutch long-term follow-up study. <u>J Viral Hepat. 2021 Jan;28(1):205-208</u> .	
Project Title:	Immunology of persistent viral infections and biomarker studies to predict	
	development of liver cancer.	
Abstract:	The innate and adaptive immune response to HBV, HCV, HEV and HIV/HCV co-infections: NK	
	and virus-specific T cells	
	Our previous studies have shown that NK cells from chronic HBV patients are functionally	
	impaired. Moreover, we and others demonstrated that the virus-specific T cell compartment in	
	chronic HBV/HCV patients is altered and not potent enough to eradicate the virus.	
	The project is aimed at characterizing the functional defect of NK cell and T cell responses in	
	patients in more detail, with special focus on the mechanisms that regulate and suppress these	
	responses. During the project peripheral blood lymphocytes and also responses in the liver	
	compartment will be assessed using flow cytometry with HBV/HCV/HIV tetramer-specific	
	multimers and functional markers. Furthermore, highly sensitive assays to determine the	
	function of NK cells and HBV/HCV-specific CD4 ⁺ and CD8 ⁺ T cells will be conducted in order to	
	identify specific markers and mechanisms that initiate and maintain the chronicity of viral	
	hepatitis infections. Besides characterization of the	
	chronic phase of infection also changes in the immune	
	response during standard-of-care and novel therapy and	
	after stopping therapy will be assessed. The studies	
	combine classical immunological studies with	
	transcriptomics/proteomics to identify biomarkers that	
	predict the response to therapy. For more information	
	see: www.viralhepatitis.nl	
	Biomarker studies in viral hepatitis and HCC	
	Hepatobiliary malignancies represent a major cause of mortality globally. The most common	
	tumors are hepatocellular carcinoma (HCC). Key factors related to the excessive mortality of	
	these tumors are the lack of reliable screening methods and the complexity of diagnosis, which	
	requires advanced imaging technology and difficult-to-access tissue. These barriers are amplified	
	by poor accessibility present in resource-limited regions, all of which leads to tumors being diagnosed at advanced stages in which curative therapy is not an ention. To every these	
	diagnosed at advanced stages in which curative therapy is not an option. To overcome these	
	barriers, we will validate immune-related markers in serum to predict HCC in South America and	
	evaluate factors associated to early HCC development.	
	This project advances the field by focusing on a unique approach to screen and diagnose tumors	
	based on serum detection of biomarkers before a tumor is visible on imaging, allowing for early	
	tumor detection in a cost-effective manner that will lead to implementation of curative	
	therapies. In addition, this project addresses modifiable risk factors for hepatobiliary tumors that	
	could be targeted for prevention.	
Requirements of	• We are looking for highly motivated, talented students with a Master degree or MD, to join our research team. The scholarship will, at least, cover subsistence allowance and an international airplane ticket.	
candidate:	• Working in the lab requires that the student has good communication skills. Therefore we have English language requirements:	
	IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs), for English speaking countries & the Netherlands: no language	
	requirements applicable.	

School/Department:	Department of Gastroenterology & Hepatology Erasmus MC	
Supervisor information:	Sonja I. Buschow, PhD	
	Email: <u>S.Buschow@erasmusmc.nl</u>	
World no 14 Gastroenterology	• Websites: Researcher - S.I. Buschow, PhD; Research group/lab - Antigen-based Immunotherapy group;	
<u>& Hepatology</u>	(Sonja Buschow LinkedIn)	
	Most important Grants:.	
	Health Holland/TKI (Dutch government) grants for the development of a peptide-based therapeutic vaccine	
	(400k€; 2017) against chronic HBV infection and its subsequent testing in a Phase I study (800k€; 2021) all in collaboration with Company ISA pharmaceuticals b.v.	
	KWF (Dutch cancer association) grants for the development of T cell therapy for liver cancer (150k€; 2020) and the	
	development of an Mass Spectrometry-based Immunopeptidomics approach to identify T cell targets (150k€;	
	2016).	
	Most important publications:	
	Jansen et al., Clin Transl Immunology. 2021 Li et al., Hepatology. 2021	
	Bouzid et al., Cancers. 2021De Beijer et al., J Virol. 2020Dou et al., J Infect Dis. 2018Worah et al., Cell Rep. 2016	
	Buschow et al., J Hepatol. 2015 Tel et al. Blood. 2013	
	Buschow et al., Traffic 2009 Van Niel et al., Immunity 2006	
Project Title:	Antigen-based Immunotherapy development for gastrointestinal & Hepatic disease	
Abstract:	Our translational research projects are aimed at finding T cell targets for antigen	
	specific immunotherapy development for different gastrointestinal and hepatic	
	diseases, including viral hepatitis and cancers.	
	For this purpose we elucidate which antigens are presented as peptides in HLA both	
	on professional antigen presenting dendritic cells (DCs) to initiate T cell responses, as	
	well as on infected or malignant cells to be targeted by effector T cells. We analyze	
	HLA-eluates by Mass spectrometry to get insight into (the regulation of) antigen	
	processing, presentation and recognition in DCs and target cells and to derive	
	effective HLA-epitopes for immunotherapy. In the lab we use various immunological	
	assays to further investigate the significance of identified epitopes, to test prototype	
	vaccines and to study regulatory mechanisms for disease specific immune responses.	
	We have already developed a therapeutic peptide based vaccine for chronic hepatitis	
	B infection that now awaits clinical testing and now aim to develop vaccines also for	
	liver cancer and other gastrointestinal malignancies. In addition we intent to improve	
	immunotherapy design and treatment regimens by researching which adjuvants or	
	immune modulatory treatments (e.g. checkpoint inhibitors) can most effectively	
	support antigen-based immunotherapy specific diseases or even patients.	
Requirements of	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills. 	
candidate:	 Master degree or MD with demonstrated experience in basic immunological and/or biochemical research techniques 	
	• Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the	
	scientific part of your scholarship proposal)	
	English language requirement: O English speaking countries & Netherlands: no requirement	
	 Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs) 	

School/Department:	Department of Gastroenterology & Hepatology Erasmus MC	
Supervisor	dr Qiuwei Abdullah Pan, <u>q.pan@erasmusmc.nl</u>	
information:	Website: <u>https://www.erasmusmc.nl/en/research/researchers/pan-q</u>	
2	Personal Grants (ongoing):	
World no 14	 Netherlands Organisation for Scientific Research, Vidi grant: € 800,000 	
Gastroenterology &	 Dutch Cancer society young investigator grant, € 549.000 	
Hepatology	Most relevant recent publications as corresponding author:	
<u>Hepatology</u>	1. LGR5 marks targetable tumor-initiating cells in mouse liver cancer. <u>Nature Communications</u> . 2020 Apr 23;11(1):1961. doi: 10.1038/s41467-020-15846-0. (IF: 15)	
	2. Cancer-Associated Fibroblasts Provide a Stromal Niche for Liver Cancer Organoids That Confers Trophic Effects and Therapy Resistance. <u>Cell Mol Gastroenterol Hepatol</u> . 2021;11(2):407-431. (IF: 9.2)	
	3. Estimating Global Prevalence of Metabolic Dysfunction-Associated Fatty Liver Disease in Overweight or Obese Adults. <u>Clinical</u> <u>Gastroenterology and Hepatology</u> . 2021 Feb 20:S1542-3565(21)00208-1. (IF: 11.4)	
	4. The biological process of lysine-tRNA charging is therapeutically targetable in liver cancer. <i>Liver International</i> . 2021 Jan;41(1):206-219. (IF: 5.8)	
	5. Dynamics of Proliferative and Quiescent Stem Cells in Liver Homeostasis and Injury. <u>Gastroenterology.</u> 2017 Oct;153(4):1133-1147. (IF: 22.7)	
	6. Unphosphorylated ISGF3 drives constitutive expression of interferon-stimulated genes to protect against viral infections. <u>Science</u> <u>Signaling</u> . 2017 Apr 25;10(476). pii: eaah4248. (IF: 8.2)	
	7. SMAD4 exerts a tumor-promoting role in hepatocellular carcinoma. <u>Oncogene</u> . 2015 Sep 24;34(39):5055-68. (IF: 9.9)	
	Publication link (about 200 in total; >20 first authorship; >100 last/corresponding authorship publications)	
	https://pubmed.ncbi.nlm.nih.gov/?term=Pan+Q%5BAU%5D+AND+%28Erasmus%29+OR+Pan%2C+Qiuwei&sort=date&size=100	
Project Title:	Understanding the biological and therapeutic implications of stem cells in liver cancer	
Abstract:	The key concept underlying the cancer stem cell (CSC) or tumor-initiating cell (TIC) theory is that	
	tumors are maintained through a hierarchical structure, in which different cell populations have	
	different functionalities in pathophysiology. The bulk of a tumor is thought to consist of CSCs/TICs	
	as well as rapidly proliferating cells. CSCs/TICs are responsible for tumor initiation, resistance to	
	conventional treatment, and distant metastasis.	
	In the liver, we previously have characterized two populations of stem cells in responding to tissue	
	injury, including the proliferative LGR5 stem cells and label-retaining quiescent stem cells. We	
	further defined that the LGR5 compartment as an important CSC population, representing a viable	
	therapeutic target for combating liver cancer.	
	Hepatitis virus infection and fatty liver disease are the main causes of liver cancer. In this project,	
	we aim to in depth understand the role of different stem cell populations in liver carcinogenesis	
	and develop potential therapeutic targeting in the context of viral hepatitis and fatty liver disease-	
	caused liver cancer.	
Poquiromonts of	We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using	
Requirements of	team work to tackle large scientific questions and thus requires a student with good communication skills.	
candidate:	Master degree or MD with demonstrated experience in basic immunological and/or biochemical research techniques	
	• Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the	
	scientific part of your scholarship proposal)	
	English language requirement: <u>English speaking countries & Netherlands:</u> no requirement	
	 English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs) 	

School/Department:	Department of Gastroenterology & Hepatology Erasmus MC
Supervisor	• dr Qiuwei Abdullah Pan, <u>g.pan@erasmusmc.nl</u>
information:	Website: <u>https://www.erasmusmc.nl/en/research/researchers/pan-q</u>
-	Personal Grants (ongoing):
World no 14	Netherlands Organisation for Scientific Research, Vidi grant: € 800,000
Gastroenterology &	• Dutch Cancer society young investigator grant, € 549,000
<u>Hepatology</u>	 Most relevant recent publications as corresponding author: 1. Potential association between COVID-19 mortality and health-care resource availability. <u>Lancet Global Health</u>. 2020
	Apr;8(4):e480. (IF: 26.8 ; Cited 530)
	2. Estimating Global Epidemiology of Low-Pathogenic Human Coronaviruses in Relation to the COVID-19 Context. Journal
	<u>of Infectious Diseases</u> . 2020 Jul 23;222(4):695-696. (IF: 5.2)
	3.Systematically mapping clinical features of infections with classical endemic human coronaviruses. <u>Clinical Infectious</u>
	<u>Diseases</u> . 2021 Aug 2;73(3):554-555. (IF: 9.1)
	4. Hepatitis E virus infection activates NLRP3 inflammasome antagonizing interferon response but therapeutically targetable. <u>Hepatology</u> . 2021 Aug 15. doi: 10.1002/hep.32114. (IF: 17.4)
	5. Cross-reactivity towards SARS-CoV-2: the potential role of low-pathogenic human coronaviruses. Lancet Microbe 2020
	Aug;1(4), e151.
	Publication link (about 200 in total; >20 first authorship; >100 last/corresponding authorship publications)
Due is at Title.	https://pubmed.ncbi.nlm.nih.qov/?term=Pan+Q%5BAU%5D+AND+%28Erasmus%29+OR+Pan%2C+Qiuwei&sort=date&size=100
Project Title:	Antiviral therapy development against human coronavirus infections
Abstract:	Coronaviruses are a large family of RNA viruses circulating among a wide range of animal species.
	Seven types of coronaviruses naturally infect humans, although all of them are thought to
	originate from animals. The three highly pathogenic coronaviruses, including MERS-CoV, SARS-
	CoV, and SARS-CoV-2, can cause severe acute respiratory diseases in humans. By contrast, the
	four genotypes of low pathogenic human coronaviruses (LPH-CoV), including OC43, HKU1, 229E
	and NL63, usually only cause mild and self-limiting respiratory tract infections. Genetically, SARS-
	CoV-2, SARS-CoV, MERS-CoV, OC43 and HKU1 are betacoronaviruses, whereas 229E and NL63 are
	alphacoronaviruses. SARS-CoV-2 is most closely related to SARS-CoV, moderately to MERS-CoV
	and is slightly distal to LPH-CoV.
	LPH-CoV, including OC43, HKU1, 229E and NL63 are endemic and have been widely circulating
	among the global population for decades. We recently have comprehensively characterized the
	clinical features of LPH-CoV and they actually can cause severe outcomes in special patient
	populations. However, there is no approved medication for treating these infections. The
	unprecedented escalation of COVID-19 pandemic has called urgency for antiviral drug
	development. In this project, we aim to understand the antiviral mechanisms and develop
	antiviral therapies against both high and low pathogenic coronaviruses as well as possible new
	coronaviruses that may emerge in the future.
Requirements of	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills.
candidate:	 Master degree or MD with demonstrated experience in basic immunological and/or biochemical research techniques
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the
	scientific part of your scholarship proposal)
	English language requirement: English spacking countries & Natherlands: no requirement
	 English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

Department of General Practice - Musculoskeletal disorders

The Department of General Practice is internationally renowned for its high-quality, innovative and multidisciplinary research on the diagnosis, prognosis and treatment of musculoskeletal disorders in primary care.

Main areas of research:

Early diagnosis, prognosis and (subgroup specific) treatment of musculoskeletal disorders, specifically:

- (1) Osteoarthritis and related disorders
- (2) Low back pain and neck/shoulder pain
- (3) Musculoskeletal disorders in the young and active individual

Why choosing for this department?

The research is led by prof.dr. BW Koes (World #4 expert on back pain) and prof.dr. SMA Bierma-Zeinstra (World #5 expert on osteoarthritis). Together with a team of assistant/associate professors (2), post-doctoral researchers (4) and over 30 PhD-students, this vibrant research group delivers high-quality research, publishes is the top international journals in the field, is well acknowledged in multiple international guideline and guideline committees, and is an active player in multiple global and multi-disciplinary research projects. Within Erasmus MC, the research group works together with departments of Orthopedics, Radiology, Medical Imaging Processing, Internal Medicine, Genetics, Sports Medicine, Epidemiology, Biomechanics, and Rheumatology to address all aspects of musculoskeletal disorders. The department works with large data sets (Rotterdam Study; CHECK, BACE, OA Trial Bank) as well as with newly collected data for diagnostic/prognostic and interventional studies.

Honors & Awards (selection)

- Editorial Board Memberships of prestigious magazines: Osteoarthritis & Cartilage (Bierma-Zeinstra; associate editor), British Journal of Sports Medicine (Middelkoop, Macri)
- Personal Awards: Clinical Research Award by the Osteoarthritis Research Society International (2015)
- Personal Grants (NWO, ERC, other)
- NWO Vidi €900K
- Collaborative Grants (NWO, Horizon2020, MSCA, other):
- NWO/ZonMw 3 mil€
- Other (inter)national funds (incl. charity) 20 mil€

Key publications of the department

Prof. BW Koes

Cochrane Database Sys Rev, 2020; 4(4):CD013581 BMJ, 2019; 367:I6273 The Lancet, 2018;391,10137 N Engl J Med, 2017;376(12):1111-1120 BMJ, 2012;344:e497 N Engl J Med, 2007;356(22):2245-56 Ann Intern Med, 2007;147(10):685-92

Prof. SMA Bierma-Zeinstra

Br J Sports Med, 2020; 54(14):822-824 Lancet, 2019; 393:1745-59 Nat Rev Rheum, 2019;15:438-448 Nat Rev Rheum, 2017;13(12):705-706 JAMA, 2017;318(12):1184 BMJ, 2017; 356:j1131 N Engl J Med, 2014;370(26):2546-7

Department of General Practice

School/Department:	'Musculoskeletal disorders' at the Department of General Practice and Department of	
	Orthopedic Surgery	
Supervisor	Prof dr SMA Bierma-Zeinstra	
information: world no 13 Surgery	Email: s.bierma-zeinstra@erasmusmc.nl	
	Website: https://www.erasmusmc.nl/en/research/groups/general-practice	
	Personal Grants:	
world no 21 Public, Environmental & Occupational Health	 Early identification and prevention of knee osteoarthritis (NWO VIDI) "Anna Prijs" (National award for excellent biomedical musculoskeletal research) Clinical Research Award of the Osteoarthritis Research Society International (OARSI) Most important publications: 	
<u>world no 32 Clinical</u> <u>Medicine</u>	- Br J Sports Med 2020; 54(14):822-824 - Nat Genetics, 2014;46(5):498-502 Lancet 2019; 393:1745-59 - JAMA, 2013;310(8):837-847 - Nat Rev Rheumatol 2019;15:438-448 - Nature Rev Rheum, 2013;9(10):630-4 - Ann Rheum Dis 2018;77:875-882 - Nat Genetics, 2011;43(2):121-6 - Nat Rev Rheum, 2017;13(12):705-706 - BMJ, 2010;341:c5688 - JAMA, 2017;318(12):1184 - JAMA, 2010;303(2):144-9 - BMJ, 2017; 356:j1131 - BMJ, 2009;339:b4074 - N Engl J Med, 2014;370(26):2546-7 - -	
Project Title:	The early diagnosis, prognosis and (subgroup specific) treatment of osteoarthritis	
Abstract:	Osteoarthritis is the most common form of rheumatic diseases. Due to the aging population and the high prevalence of overweight and obesity, the prevalence of osteoarthritis is rising. In the Netherlands, osteoarthritis is expected to be the most prevalent disease by 2040. The majority of patients with osteoarthritis are treated in primary care and orthopedic practice. Early diagnosis, identification of high-risk groups, and surrogate outcomes in early OA can help optimizing treatment for patients with osteoarthritis, or even prevention. As there is no cure for osteoarthritis, current treatment focusses on symptomatic relief. On average, treatment effects of guideline recommended treatments for osteoarthritis provide small to moderate improvements in pain and function. Nevertheless, subgroups of patient with osteoarthritis do respond strongly to certain types of interventions and should hence be identified for optimal treatments effect. Within this internationally renowned research group, multiple research projects on the epidemiology and (subgroup specific) treatment of osteoarthritis in primary care are available for highly motivated junior researchers.	
Requirements of candidate:	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using teamwork to tackle large scientific questions and thus requires a student with good communication skills. Master degree or MD Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) English language requirement: English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs) 	

Department of General Practice

School/Department:	'Musculoskeletal disorders' at the Department of General Practice
Supervisor	Prof dr BW Koes
information:	Email: <u>b.koes@erasmusmc.nl</u>
	Website: <u>https://www.erasmusmc.nl/en/research/groups/general-practice</u>
world no 21 Public,	Personal Grants:
Environmental & Occupational Health	- Advise and medical treatment of acute low back pain in primary care (NWO)
	- Medical treatment of sciatica in primary care (NWO)
world no 32 Clinical	Most important publications:
<u>Medicine</u>	- Cochrane Database Sys Rev, 2020; 4(4):CD013581
	- BMJ, 2019; 367:l6273
	- The Lancet, 2018;391,10137
	- N Engl J Med, 2017;376(12):1111-1120
	- BMJ, 2012;344:e497
	- N Engl J Med, 2007;356(22):2245-56
	- Ann Intern Med, 2007;147(10):685-92
Project Title:	Diagnosis and prognosis of musculoskeletal disorders
Abstract:	 Musculoskeletal disorders occur very frequently in primary care. The etiology, diagnosis and prognosis are often unknown, which hampers adequate management of patients presenting with these disorders in primary care. Our department is one of the international key-players in the field of musculoskeletal disorders in primary care. We are involved in a large number of cohort studies and clinical trials evaluating risk factors, the value of diagnostic- and therapeutic interventions, as well as studying the prognosis (and its determinants) of the most common musculoskeletal disorders presenting in primary care. This includes studies on low back pain, sciatica, neck and shoulder pain, knee pain (patellofemoral pain syndrome), ankle distortions, and osteoarthritis. We also study musculoskeletal disorders and sport injuries among the young and active individuals. Next to original research, the department is also active in writing systematic reviews and metaanalysis on these topics. The PhD-candidate will be active with (secondary) data-analysis, writing original research papers and systematic reviews within the field of musculoskeletal disorders in primary care.
Requirements of candidate:	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills. Master degree or MD Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) English language requirement: English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

Department of Hospital Pharmacy

Department:	Department of Hospital Pharmacy, Erasmus MC
Supervisor	Prof. dr.P.H.M. (Hugo) van der Kuy, Prof. dr. K.M. (Karel) Allegaert, Prof. dr. B.C.P. (Birgit) Koch,
information:	Associate prof. dr. L.E. (Loes) Visser
	Email research coordinator: <u>e.e.m.vankampen@erasmusmc.nl</u>
<u>world no 39 in</u>	Website: <u>https://www.erasmusmc.nl/en/research/departments/pharmacy</u>
Pharmacology &	Grants: Several national grants, IMI and the Combacte grant from European Union.
<u>Toxicology</u>	Most important publications:
	Abdulla, Alan et al. "Failure of Target Attainment of Beta-Lactam Antibiotics in Critically III Patients and Associated Risk Factors: A Two-Center Prospective Study (Expat)." Critical Care 24, no. 1 (2020/09/15 2020): 558. <u>https://doi.org/10.1186/s13054-020-03272-z</u> .
	Atrafi, Florence et al. "Intratumoral Comparison of Nanoparticle Entrapped Docetaxel (Cpc634) with Conventional Docetaxel in Patients with Solid Tumors." Clinical Cancer Research 26, no. 14 (2020): 3537. <u>https://doi.org/10.1158/1078-0432.Ccr-20-0008</u> . Francke, M. I. et al. "Monitoring the Tacrolimus Concentration in Peripheral Blood Mononuclear Cells of Kidney Transplant Recipients." Br J Clin Pharmacol (Oct 6 2020).
	Kloosterboer, S. M. et al. "Risperidone Plasma Concentrations Are Associated with Side Effects and Effectiveness in Children and Adolescents with Autism Spectrum Disorder." Br J Clin Pharmacol (Jul 9 2020).
	Sablerolles, R. S. G., et al. "Covid Medication (Comet) Study: Protocol for a Cohort Study." Eur J Hosp Pharm 27, no. 4 (Jul 2020): 191-93.
-	Van den Anker, J. N., et al. "Approaches to Dose Finding in Neonates, Illustrating the Variability between Neonatal Drug Development Programs." Pharmaceutics 12, no. 7 (Jul 20 2020).
Project Title:	PhD-projects in the hospital pharmacy, Erasmus MC
Abstract:	Within our pharmacy, the goal is to individualize and optimize patient drug therapy. To achieve this our
	research is built on three research lines:
	1. <u>Medication optimization and safety</u>
	Research focused on the optimization of pharmacotherapy in primary care and in secondary or tertiary
	care settings. This domain also works on prevention of (re-)hospitalizations by optimizing
	pharmacotherapy. Within this research line, there is an epidemiological track.
	Head of department, prof. dr. P.H.M. (Hugo) van der Kuy, associate prof. dr. J. (Jorie) Vermissen,
	associate prof. dr. L.E. (Loes) Visser
	2. <u>Model-based dosing</u>
	No two patients are identical, so individual drug dosing can lead to better treatment. The focus is on
	pharmacokinetics (PK) and pharmacodynamics (PD), therapeutic drug monitoring (TDM), and their
	implementation in clinical practice. By the use of PK/PD models we establish the relation between
	drug dosage, drug concentration and drug effect and we implement the outcomes of our research in clinical practice. Principal investigator, associate professor, <u>dr. B.C.P. (Birgit) Koch</u> .
	3. <u>Pediatric and perinatal pharmacology</u>
	This research line includes different topics; prescribing to children, advanced therapy medicinal
	product (ATMP), oncology and radio-pharmacy. For children PK/PD modeling is a good way to
	achieve safe prescriptions of (off-label) drugs in neonatal intensive care. With the opening of our
	ATMP facility we are combining fundamental research and clinical practice. Furthermore we are
	innovative in the field of radio-pharmacy by labeling specific tracers. Upon that we are planning
	trials with 3D-printed tablets to optimize individual dosing. Team, prof. dr. K.M. (Karel) Allegaert, dr.
	R.B. (Robert) Flint , dr. E.J. Ruijgrok and dr. S.L.W. (Stijn) Koolen.
	Within these research lines, we also investigate education; for example the most effective teaching
	tools for medical students. Principal investigator, assistant professor, dr. F. (Floor) van Rosse.
	Further information: https://www.erasmusmc.nl/en/research/departments/pharmacy
Requirements of	We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using
candidate:	team work to tackle large scientific questions and thus requires a student with good communication skills. The candidate
	should have great interest in the field of pharmacy, medication optimization, pharmacometrics, modelling and/or
	 pediatric pharmacology. Master degree or MD in pharmacy, medicine, biomedical or biopharmaceutical sciences
	 Master degree or MD, in pharmacy, medicine, biomedical or biopharmaceutical sciences. Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we will help with the
	scientific part of your scholarship proposal)
	 English language requirement:
	• English speaking countries & Netherlands: no requirement
	 Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

Department of Immunology

The mission of the Department of Immunology at Erasmus University Medical Center is to perform cutting edge and outstanding fundamental, translational and clinical research, provide excellent teaching in Immunology and support patient care with high quality immunological diagnostic services. Research in the department of Immunology spans molecular to clinical immunology and includes the development and function of innate and adaptive immunity, autoimmunity and inflammation, immune deficiencies, immunity to pathogens and tumors, neuroimmunology, computational biology in immunology and lymphoid malignancies.



The department of Immunology and its faculty have a long history of excellent training of PhD students in an intellectually stimulating and culturally diverse environment. The department of Immunology has state-of-the-art research facility, including bioinformatics, and provides an outstanding environment for PhD student training. Faculty of the department have extensive national and international collaborations, and a seminar series that provide excellent opportunities for students to network. Further information on the department, individual faculty and programs can be found at: https://www.erasmusmc.nl/immunologie/?lang=en.

Key publications 2020-21 by PI's of the Dept. of Immunology

- Assmann, Jorn L.J.C. et al. 2021. "TRB Sequences Targeting ORF1a/b Are Associated with Disease Severity in Hospitalized COVID-19 Patients." Journal of Leukocyte Biology. (September 15, 2021).
- Erkeland, Stefan J et al. 2021. "The MiR-200c/141-ZEB2-TGFβ Axis Is Aberrant in Human T-Cell Prolymphocytic Leukemia." Heamatologica.
- Meijers, Ruud W.J. et al. 2020. "Responsiveness of Chronic Lymphocytic Leukemia Cells to B-Cell Receptor Stimulation Is Associated with Low Expression of Regulatory Molecules of the Nuclear Factor-KB Pathway." *Haematologica* 105(1): 182. (September 15, 2021).
- Mueller, Yvonne M et al. 2021. "Immunophenotyping and Machine Learning Identify Distinct Immunotypes That Predict COVID-19 Clinical Severity." medRxiv: 2021.05.07.21256531. (May 18, 2021).
- Orme, Michelle E. et al. 2021. "Systematic Review of Anti-DsDNA Testing for Systemic Lupus Erythematosus: A Meta-Analysis of the Diagnostic Test Specificity of an Anti-DsDNA Fluorescence Enzyme Immunoassay." Autoimmunity Reviews: 102943. (September 15, 2021).
- van Riet, Job et al. 2021. "The Genomic Landscape of 85 Advanced Neuroendocrine Neoplasms Reveals Subtype-Heterogeneity and Potential Therapeutic Targets." *Nature Communications* 12(1): 1–14. (July 29, 2021).
- Schrijver, Benjamin et al. 2020. "Inverse Correlation between Serum Complement Component C1q Levels and Whole Blood Type-1 Interferon Signature in Active Tuberculosis and QuantiFERON-Positive Uveitis: Implications for Diagnosis." *Clinical & Translational Immunology* 9(10): e1196. (September 15, 2021).
- van der Velden, Vincent H. J. et al. 2021. "Potential and Pitfalls of Whole Transcriptome-Based Immunogenetic Marker Identification in Acute Lymphoblastic Leukemia; a EuroMRD and EuroClonality-NGS Working Group Study." *Leukemia 2021 35:3* 35(3): 924–28. (September 15, 2021).
- Talarico, Rosaria et al. 2021 "The impact of COVID-19 on rare and complex connective tissue diseases: the experience of ERN ReCONNET". Nature Reviews Rheumatology 2021 17(3):177-84
- Tyler, Paul M. et al. 2021. "Human autoinflammatory disease reveals ELF4 as a transcriptional regulator of inflammation". Nature Immunology 2021 22(9): 1118-26
- Zhao, Manzhi et al. 2020. "Rapid in Vitro Generation of Bona Fide Exhausted CD8+ T Cells Is Accompanied by Tcf7 Promotor Methylation" ed. Annette Oxenius. *PLOS Pathogens* 16(6): e1008555. (November 24, 2020).

Editorial Board Memberships:

Associate Editor, Frontiers in Immunology (Katsikis); Review Editor, Frontiers in Genetics (Katsikis); Editorial Board Member in Cells and in BioMedInformatics (van de Werken), Section Editor, Journal of Immunology (Katsikis till 2014)

The department has a track record of external funding via grant support. Selected grants mentioned:

Horizon2020 (Drexhage), NWO Vidi (van Luijn; van der Burg) and Aspasia (van der Burg), NWO-VENI award, KWF-fellowship and cancer research grants (Erkeland), Worldwide Cancer Research Grant and NIH (Katsikis), DDHF (van de Werken), ReumaFonds (Versnel), Prinses Beatrix Spierfonds and Horizon2020 (Jacobs) and pharma industry (Langerak, van der Velden, van Hagen).

Department of Immunology

School/Department:	Department of Immunology, Erasmus MC	
Supervisor information:	•Prof dr. P. Martin van Hagen; <u>p.m.vanhagen@erasmusmc.nl</u>	
world no 31 in Immunology	 Grants: IPAD trial: Influencing Progression of Airway Disease in patients with Primary Antibody DeficiencyGenetics first in Primary Immune Deficiency, Netherlands Organisation for Health Research and Development, 2019 PIPGEN Project 7 : The role of PI3K neurodevelopmental disorders: Marie Sklodowska-Curie Grant , EU Horizon 2020, 2020 Moodstratiification: EU Horizon 2020, 2018 Co-supervisor: Dr. Virgil A.S.H. Dalm Co-supervisor: Dr. Layal Chaker Secondary affiliation dr. Chaker: Harvard T.H. Chan School of Public Health Most important publications of supervisors: J Allergy Clin Immunol. 2016, PMID: 31268374 Lancet, 2017, PMID: 28336049 	
	Nature Communications., 2020, PMID: 32769997 Nature Immunology , 2021 PMID: 34326534	
Project Title:	J Clin Immunol., 2021, PMID: 34505230 Nat Rev Rheumatoly, 2021, PMID: 33408338 Deciphering the genomic and epi-genomic landscape of immunoglobulins	
Abstract:	Immunoglobulins (Igs) have a central role in the immune response by specifically recognizing	
	 and binding to particular antigens, such as bacteria or viruses, and aiding in their abolishment. The antibody immune response is highly complex and has recently gained general interest during the COVID-19 pandemic. Also, Igs, as well as the immune system in general, have been attributed a critical role in inflammation and inflammaging, potentially providing a viable target for age-related diseases such as cardiovascular disease (CVD). While certain environmental aspects influencing fluctuations and differences in serum levels of Igs have been uncovered, there is still little to no information on the genomic landscape involved in this process. Furthermore, differences in methylation, a process that can change DNA activity without changing its sequence, that may lead to differences between Igs and Ig response in the population, has never been study, but may be crucial. Unravelling essential genetic variations is pivotal for several outstanding issues including antibody responses to infections or vaccinations as well as clinically relevant diseases (e.g. immunodeficiency disorders). With this project we aim to decipher the genomic and epigenomic (methylation) landscape of immunoglobulins. We will use genome-wide (GWAS) approaches to identify novel genetic variations responsible for immunoglobulin levels and responses with in the general population. Investigating whether methylation pattern differences in the general population are associated with differences for immunoglobulin levels and response through a so-called Epigenome-wide association study (EWAS) Construct polygenic risk scores to investigate potential causal association with inflammaging and inflammation-associated diseases, such as CVD and cancer. Utilize Mendelian Randomization approaches for studying causality between immunoglobulins and age-related diseases. We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in 	
Requirements of candidate:	 We are footing for a highly indivated, hardworking student to join out very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills. Master degree or MD with a background in statistical programming, preferably R Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) English language requirement: English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs) We offer you: Overhead and material costs Fees for relevant coursework and conferences 	

Department of Immunology

School/Department:	Department of Immunology Erasmus MC
Supervisor information:	Prof dr. Anton W Langerak (supervisor)
	• Dr. Harmen JG van de Werken & Dr. Marco WJ Schreurs (co-supervisors)
world no 31 in Immunology	Email: <u>a.langerak@erasmusmc.nl</u> and/or <u>h.vandewerken@erasmusmc.nl</u> and/or <u>m.schreurs@erasmusmc.nl</u> Website: <u>Anton Langerak</u> and <u>Harmen van de Werken</u> & <u>II</u> and <u>Marco Schreurs</u> Personal Grants:
	1. DDHF CCBC (2018)
	2. EU-TRANSCAN NOVEL (2019)
	Most important recent relevant publications:
	 van de Werken, H. J. G.*, van Riet, J.*,, Mostert, B. The genomic landscape of 85 advanced neuroendocrine neoplasms reveals subtype-heterogeneity and potential therapeutic targets. <i>Nat. Commun.</i> 12, 1–14 (2021).
	 Assmann JLJC*, Kolijn PM*, Schrijver B*, Langerak AW. TRB sequences targeting ORF1a/b are associated with disease severity in hospitalized COVID-19 patients. J Leukoc Biol. 2021. Epub ahead of print.
	- van Riet, J.,, van de Werken, H. J. G. SNPitty: An Intuitive Web Application for Interactive B-Allele Frequency and Copy
	Number Visualization of Next-Generation Sequencing Data. <i>J. Mol. Diagnostics</i> 20, 166–176 (2018).
	 van de Werken, H. J. G.,, Joffe, B. Small chromosomal regions position themselves autonomously according to their chromatin class. <i>Genome Res.</i> 27, 922–933 (2017).
	 van de Werken, H. J. G.*, Landan, G*.,, de Laat, W. Robust 4C-seq data analysis to screen for regulatory DNA interactions.
	Nat. Methods 9, 969–972 (2012)
Project Title:	Precision medicine in an immune disease and cancer context using Machine learning
	and Artificial intelligence
Abstract:	Machine Learning (ML) and Artificial Intelligence (AI) are key to better predict clinical outcome with
	highly complex clinical and molecular data sets. Moreover, these sophisticated methods can be applied
	to develop new algorithms and visualization tools to better understand basic cellular and molecular
	principles. In this project we aim to improve our biological understanding, diagnostic tools and
	response to therapy through ML and AI using different context-dependent -omics data sets in three
	subprojects:
	1. We will deeply interrogate whole transcriptome data to understand transcription and aberrant
	splicing in cancer. We will develop new algorithms ⁵ and visualization tools ³ and integrate whole
	genome data and chromosome conformation data when necessary ^{1,4} . This can lead to many novel
	insights in cancer development and potential new therapies in this devastating disease.
	2. We will use immune receptor repertoire ("immunome") data from lymphoproliferative disease to
	identify context-dependent profiles of immune cells ² . These profiles can support precision medicine
	through 1) definition of benign and malignant immune cell clones (diagnostics/prognostics) 2)
	traceability of clones upon therapy (monitoring), and 3) identification of disease-specific patterns to guide therapeutic decision making (theranostics). Examples of the impact of immunome analysis in a
	broader context include: Stereotyped BCR subsets in chronic leukemia with different prognostics,
	minimal disease monitoring, eligibility for immune therapy, TCR profiles with disease impact in cancer
	but also infectious disease, e.g. COVID-19.
	3 . We aim to improve allergy diagnostics based on the IgE profile of allergic individuals. The newly
	developed Allergy Explorer (ALEX) allows the acquisition of an IgE profile comprising 282 allergen
	extracts and components. The major challenge is the correct and clinically useful interpretation of such
	extensive IgE profiles, including reactivity of variable clinical implication. AI may support the clinician in
	the interpretation of the IgE profiles in combination with clinical signs and symptoms, and other clinical
	and demographic patient characteristics.
	Based on these projects we hope to show that ML and AI supported clinical decision making as such
	may significantly benefit future treatment of cancer and immunological disease at a personal level
	(Precision Medicine).
Requirements of	• We are looking for a candidate with strong analytical and problem-solving skills, being highly motivated and having
candidate:	excellent communication and writing skills and being able to work independently. A background in immunology and/or cancer biology is of significant added value.
	 Master's degree in bioinformatics, computational biology, statistics, or a related field.
	 The candidate should have demonstrated excellent scientific writing and software engineering skills in R and Python
	or Perl.
	• Scholarship that will, at least, cover subsistence allowance and international airplane ticket (we could help with the
	scientific part of your scholarship proposal)
	English language requirements
	 English language requirement: English speaking countries & Netherlands: no requirement

Department of Internal Medicine – Calcium & Bone Metabolism

Why would you do scientific research on bone?

Contrary to general belief, the skeleton is a highly dynamic organ where many energy demanding processes take place, such as life-long bone remodeling, stem cell renewal, hematopoiesis and mineral homeostasis. Therefore, bone plays a central role in a wide variety of diseases affecting millions of people world-wide.

Our international team is working on 3 main research lines: 1) Bone regeneration: We aim to characterize the mechanisms behind bone cell differentiation and underlying bone formation and degradation to gain insight into diseases where bone formation is not well controlled (osteoporosis, craniosynostosis) or during fracture healing. 2) Bone metastases: We study the complex interactions between bone metastatic cancer cells and osteoblasts to identify new therapeutic approaches in bone metastases and potentially diagnostic profiles. 3) Rare bone diseases: We investigate the molecular mechanisms of rare, monogenic human diseases of disturbed bone and mineral metabolism as well as candidate bone anabolic genes derived from large population-based genetic studies.

Group of Calcium & Bone metabolism: we have trained over 25 PhD students and have published around 250 papers. Our team has been involved in numerous (inter)national collaborations/grants, and we list a few European grants to give you an impression:

- FP6: GEFOS, NucSys (Marie Curie RTN)
- FP7: GENOMOS, PEOPLE IRSES network INTERBONE, BioInspire
- Horizon2020: MCSA-RISE

Publications:

- Lodberg A et al. A follistatin-based molecule increases muscle and bone mass without affecting the red blood cell count in mice. FASEB J. 2019;33(5):6001-6010
- Mumtaz N et al. Zika virus infection perturbs osteoblast function. Sci Rep. 2018;8(1):16975
- Brum A et al. Mucin 1 (Muc1) deficiency in female mice leads to temporal skeletal changes during aging. JBMR Plus. 2018;2(6):341-350
- Baroncelli M et al. Human osteoblast-derived extracellular matrix with high homology to bone proteome is osteopromotive. Tissue Eng Part A. 2018;24(17-18):1377-1389
- Koek N et al. Osteoclastogenic capacity of peripheral blood mononuclear cells is not different between women with and without osteoporosis. Bone. 2017;95:108-114
- Morhayim J et al. Osteoblasts secrete miRNA-containing extracellular vesicles that enhance expansion of human umbilical cord blood cells. Sci Rep. 2016;6:32034
- Brum A et al. Connectivity Map-based discovery of parbendazole reveals targetable human osteogenic pathway. Proc Natl Acad Sci U S A. 2015;112(41):12711-6

Contact information: Dr. Bram CJ van der Eerden, <u>b.vandereerden@erasmusmc.nl</u>, +31(10)7032841, @eerden1970, Skype: bramvandereerden; website: <u>https://publons.com/researcher/2698444/bram-cj-van-der-eerden/</u>

School/Department:	Department of Internal Medicine-Calcium and bone metabolism, Erasmus MC	
Supervisor	Bram C.J. van der Eerden, PhD; <u>b.vandereerden@erasmusmc.nl</u>	
information:	Website:	
	- https://www.erasmusmc.nl/en/research/researchers/eerden-bram-van-der	
world no 29 Endocrinology	- https://publons.com/researcher/2698444/bram-cj-van-der-eerden/	
<u>& Metabolism</u>	Personal grants:	
	- 2018-2022: Health~Holland, TKI,	
	- 2016-2020: Horizon2020-MCSA-RISE-2015	
	- 2012-2016: FP7-PEOPLE-2011-IRSES	
	 Most important publications (Total publications, 96; H-index, 26) 	
	- Brent et al., <u>Bone. 2021</u> ; 142: 115692	
	- Van Hengel et al., <u>Mater Today Bio. 2020</u> ; 7: 100060	
	- Fecher-Trost et al. J Bone Miner Res. 2019;34(4):699-710	
	- Lodberg et al. <u>FASEB J. 2019</u> ;33(5):6001-6010	
	- Brum et al. JBMR Plus. 2018;2(6):341-350	
	 Mumtaz et al. <u>Sci Rep. 2018</u>;8(1):16975 Vermeij et al. <u>Nature. 2016</u>;537(7620):427-431 	
	 Zambetti et al., <u>Cell Stem Cell</u>, 2016; 19(5): 613-627 	
	 Brum et al. Proc Natl Acad Sci U S A. 2015;112(41):12711-6 	
Project Title:	Integrative approach to study bone regeneration	
-	Contrary to common belief, bone is a highly dynamic and vital organ with a multitude of events	
Abstract:		
	taking place, such as continuous bone remodeling, stem cell renewal, hematopoiesis, mineral	
	homeostasis, etc. Osteoporosis, in which often several of these processes are affected, is the	
	most common skeletal disorder, affecting many millions of patients globally. As a consequence,	
	every 3 seconds an individual suffers from a fracture worldwide, of which 10% does not heal well	
	(non-union fractures). Given its complexity and multitude of cell types involved, it is difficult to	
	study specific processes taking place in the regenerating skeleton <i>in vivo</i> .	
	Within the laboratory of Calcium and bone metabolism, we therefore use a multidisciplinary	
	approach to identify new factors and mechanisms involved in bone formation and bone	
	regeneration. We study bone formation and healing in human bone cell models by manipulating	
	genes of interest and the consequences for mesenchymal stromal cell-derived osteogenesis and	
	adipogenesis and the effects on other cell types in the bone marrow niche including endothelial	
	cells. Promising new candidates are also being scrutinized in <i>in vivo</i> osteoporosis and bone	
	fracture/regeneration models. Among the currently employed state-of-the-art methodologies,	
	we use organ-on-chip (OoC) microfluidics to study cell-cell interaction under physiological cues,	
	CrispR-Cas9-mediated gene editing but also biomaterial sciences and 3D (bio)printing.	
	By studying a combination of bone formation, angiogenesis, 3D-printed scaffolds and newly	
	discovered genes/compounds, we obtain insights into novel physiologically relevant and	
	targetable processes in bone metabolism and provide a better understanding towards	
	therapeutic approaches to improve bone regeneration and shorten the societal and financial	
	burden associated with fractures.	
	The qualified candidate will work within international teams of scientists in an interdisciplinary	
	setting, and will receive both theoretical training and hands-on training in a large range of	
	cutting-edge techniques. PhD students are supported by a supervision committee, participate in	
	scientific and professional skills courses, attend international conferences and receive career	
	development support.	
Requirements of	Background: Cell biology, molecular biology, biomedical, creative, punctual, enthusiastic, communicative	
candidate:	 Master degree or MD, animal experimentation permit is preferred. 	
cunuluite:	• Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal)	
	English language requirement:	
	• English speaking countries & Netherlands: no requirement	
	O Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)	

Dept of Internal Medicine – Calcium & Bone Metabolism

School/Department:	Department of Internal Medicine-Calcium and bone metabolism, Erasmus MC
Supervisor information:	Dr. Marjolein van Driel, Prof. Dr. Hans van Leeuwen
	m.vandriel@erasmusmc.nl, j.vanleeuwen@erasmusmc.nl
world no 29 Endocrinology &	https://www.erasmusmc.nl/en/research/groups/laboratory-for-calcium-and-bone-metabolism
<u>Metabolism</u>	Recent publications:
	J Cell Physiol. 2020 May;235(5):4865-4877. doi: 10.1002/jcp.29365 FASEB J. 2020 Apr;34(4):5435-5452. doi: 10.1096/fj.201902610R
	Front Bioeng Biotechnol. 2019 Mar 1;7:38. doi: 10.3389/fbioe.2019.00038.
	FASEB J. 2019 May;33(5):6001-6010 J Cell Physiol. 2019 Mar;234(3):2984-2996
	Eur J Immunol. 2018 Feb;48(2):220-229
	Tissue Eng Part A. 2018 24(3-4):207-218
	Adv Healthc Mater. 2018 e1800507. 2018 doi: 10.1002/adhm.201800507 Bone 2018 117:70-8
	J Bone Miner Res. 2018 33(4):606-620
	J Cell Physiol. 2018 doi: 10.1002/jcp.27116
	Tissue Eng Part A. 2018 24(17-18):1377-1389 J Cell Physiol. 2018 233(1):387-395
	J Cell Physiol. 2018 233(6):4895-4906
	J Cell Physiol. 2018 233(2):1424-1433 Mol Cell Endocrinol. 2017 453:46-51
	Biochim Biophys Acta. 2017 1864(7):1133-1141
	Stem Cell Reports. 2017 Apr 11;8(4):947-960
Project Title:	Dormant cells (cancer stem cells) in bone metastases
Abstract:	The special milieu of the bone environment provides a fertile soil for many cancers to
	metastasize to. But especially for patients with breast or prostate tumors, metastatic cells
	preferentially go to the bone. The consequences of bone metastases are devastating and
	patients die because of complications to the bone. Despite the discovery of many factors
	involved, no cure has yet been found for bone metastases. The metastatic process is
	determined by highly specific interactions between disseminating cancer cells and the bone
	microenvironment.
	Recent research in our lab focuses on the role of the osteoblasts (bone forming cells) in
	metastatic growth. We developed co-culture models of osteoblasts and different types of
	metastatic prostate cancer cells (bone or non-bone derived). Only bone derived metastatic
	cancer cells can survive and grow in bone by impairing osteoblast differentiation and so keep
	osteoblasts in a tumor cell growth stimulatory stage: a vicious circle.
	When cancer cells metastasize to the bone, they can stay dormant for years in the bone before
	colonization and expansion takes place. These dormant cells are thought to be the cancer stem
	cells.
	Finding markers to trace these dormant cells and exploring the mechanisms that trigger
	these dormant cells to start proliferating in the bone environment are the main goals of the
	current PhD project.
	By performing co-culture models of differentiating osteoblasts and surviving (dormant)
	metastatic prostate cancer cells, we obtained gene profiles (micro-array) that specifically
	characterize these dormant cancer cells. These will be the basis to further discover new
	(protein) markers. Functional studies will focus on re-activation of dormant cells and studies to
	unravel the factors in the bone that trigger re-activation of dormant cancer cells.
	We will make use of GFP transduced human metastatic prostate cancer cells to be able to
	distinguish them from human osteoblasts.
	The obtained knowledge will be used to develop new therapies for bone metastases
Requirements of	 Background: Cell biology, molecular biology, interest in cancer research, creative, punctual, enthusiastic, communicative
candidate:	Master degree or MD
	• Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the
	 scientific part of your scholarship proposal) English language requirement:
	English speaking countries & Netherlands: no requirement

Dept of Internal Medicine – Cardiovascular Pharmacology

School/Department:	Department of Internal Medicine-Cardiovascular Pharmacology, Erasmus MC
Supervisor information:	Prof. Dr. Antoinette Maassen van den Brink
	Email: a.vanharen-maassenvandenbrink@erasmusmc.nl
world no 39 Pharmacology &	Website: <u>https://pharma.erasmusmc.nl/migraine.html</u>
<u>Toxicology</u>	Grants: Dutch Research Council: Veni (2004), Vidi (2011), Vici (2020)
	- Conacyt: several grants (3x postdoc, 2x PhD student)
	- Secretaría de Eduacación, Ciencia, Tecnología e Innovación. Mexico City (1x postdoc)
	- Dutch Heart Foundation
	- Dutch Brain Foundation
	- Berlin Institute of Health
	Most important publications:
	1. Van Casteren, D.S., Kurth, T., Danser, A.H.J., Terwindt, G.M., MaassenVanDenBrink, A. (2021). Sex
	differences in response to triptans: A systematic review and meta-analysis. Neurology, 96:162-170.
	2. MaassenVanDenBrink, A., Reekers, M., Bax, W.A., Ferrari, M.D., Saxena, P.R. (1998). Coronary side effect
	potential of current and prospective antimigraine drugs. <u>Circulation, 98:25 30.</u>
	3. MaassenVanDenBrink, A., Meijer, J., Villalón, C.M., Ferrari, M.D. (2016). Wiping out CGRP - potential
	 cardiovascular risks. <u>Trends in Pharmacological Sciences, 37:779-88</u>. 4. De Vries, T., MaassenVanDenBrink, A. (2019). Monoclonal antibody targeting CGRP in difficult-to-treat
	migraine. <u>Nature Reviews Neurology</u> , 15:688-689.
	5. Al-Hassany, L., MaassenVanDenBrink, A. (2020). Targeting CGRP in migraine: a matter of choice and dose.
	Lancet Neurol, 19:712-713.
	6. Mulder, I.A., Li, M., de Vries, T., Qin, T., Yanagisawa, T., Sugimoto, K., van den Bogaerdt, A., Danser, A.H.J.,
	Wermer, M.J.H., van den Maagdenberg, A.M.J.M., MaassenVanDenBrink, A., Ferrari, M.D., Ayata, C.
	(2020). Anti-migraine CGRP receptor antagonists worsen cerebral ischemic outcome in mice, <u>Ann Neurol</u> , 88:771-784.
	7. MaassenVanDenBrink, A., Meijer, J., Villalón, C.M., Ferrari, M.D. (2016). Wiping out CGRP - potential
	cardiovascular risks. Trends in Pharmacological Sciences, 37:779-88.
Project Title:	Migraine: the role of CGRP and cardiovascular safety of CGRP (receptor) blockade
Abstract:	Background: Migraine is a highly disabling and prevalent disorder, occurring 2-3 times more
	often in females than in males. A novel class of antimigraine drugs consists of antibodies
	against Calcitonin Gene-Related Peptide (CGRP) or its receptor. While blocking CGRP may be a
	big advantage for migraine patients without a good response to current therapies, the
	potential risks of 'wiping out' the vasodilator CGRP, which is thought to have a rescue
	function in case of threat of ischemia, should be well studied. Further, the role of CGRP may
	be different in male and female migraine patients, which is relevant in view of the
	I predominance of migraine in females
	predominance of migraine in females. Project description: The current PhD project will focus on the (neuro)vascular role of CGRP
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	 Project description: The current PhD project will focus on the (neuro)vascular role of CGRP, with a special emphasis on the role of sex hormones on the CGRP-ergic system. We will use animal in vivo models as well as human blood vessels in vitro. Depending on the interest of the PhD student, also human in vivo and/or epidemiological studies could be part of this project. Expected result: A typical Dutch PhD thesis, containing multiple published papers in top pharmacological or neurological journals. The PhD student will work with an extensive team of basic scientists, clinicians, and technicians, allowing him/her to cover both preclinical and
	 Project description: The current PhD project will focus on the (neuro)vascular role of CGRP, with a special emphasis on the role of sex hormones on the CGRP-ergic system. We will use animal in vivo models as well as human blood vessels in vitro. Depending on the interest of the PhD student, also human in vivo and/or epidemiological studies could be part of this project. Expected result: A typical Dutch PhD thesis, containing multiple published papers in top pharmacological or neurological journals. The PhD student will work with an extensive team of basic scientists, clinicians, and technicians, allowing him/her to cover both preclinical and clinical research.
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• •	 Project description: The current PhD project will focus on the (neuro)vascular role of CGRP, with a special emphasis on the role of sex hormones on the CGRP-ergic system. We will use animal in vivo models as well as human blood vessels in vitro. Depending on the interest of the PhD student, also human in vivo and/or epidemiological studies could be part of this project. Expected result: A typical Dutch PhD thesis, containing multiple published papers in top pharmacological or neurological journals. The PhD student will work with an extensive team of basic scientists, clinicians, and technicians, allowing him/her to cover both preclinical and clinical research. PhD student profile: Ideally, the student has a solid background in physiology and pharmacology, and some experience with animal research, biochemistry and molecular biology. He/she does not need to be a clinician. We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills. Master degree or MD Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the

School/Department:	Department of Internal Medicine-Genetics Lab & Population Genomics, Erasmus MC
Supervisor information:	Prof dr. M.C. (Carola) Zillikens; Email: <u>m.c.zillikens@erasmusmc.nl</u> Websites: <u>http://qlimdna.org/; https://www.erasmusmc.nl/en/research/groups/genetic-laboratory-of-internal-medicine;</u>
world no 29 Endocrinology & Metabolism	 https://www.erasmusmc.nl/en/research/researchers/zillikens-carola; https://www.erasmusmc.nl/en/research/groups/laboratory-for-calcium-and-bone-metabolism Grants: Several grants from Dutch and Australian Government and private foundations Most important publications: Waqas K, Chen J, et al. J Bone Miner Res. 2020 May 28. doi: 10.1002/jbmr.4096. van den Beld AW., Lancet Diabetes Endocrinol. 2018 Aug;6(8):647-658 Jiang X, et al. Nat Commun. 2018 Jan 17;9(1):260. Zillikens MC*, et al Nature Commun 2017 Jul 19;8(1):80. Erratum in: Nat Commun. 2017 Nov 7;8(1):1414. Zheng HF, et al. Nature. 2015 Oct 1;526(7571):112-7 Locke AE, et al. Nature. 2015 Feb 12;518(7538):197-206. Shungin D, et al. Nature. 2015 Feb 12;518(7538):197-96. van Dijk FS*, Zillikens MC*, et al. N Engl J Med. 2013 Oct 17;369(16):1529-36. Zhu H, et al. Cell. 2011 Sep 30;147(1):81-94.
Project Title:	10. <u>Kilpelainen TO, et al. Nat Genet. 2011 Aug;43(8):753-60</u> Advanced glycation end products in relation to ageing & age-related diseases
Abstract:	Advanced glycation end products (AGEs) are heterogeneous glycated products that accumulate in the body over lifetime as part of normal ageing but increased under certain conditions. It is becoming more and more clear that they are involved in age-related related diseases as evidence from population studies and wet-lab studies accumulates (Singh et al. 2012). AGEs (e.g. glucospane, pentosidine and carboxymethyllysine) are produced after glycation of protein amino acid residues, lipids or nucleic acids and sometimes through oxidation without enzymatic catalysis (Vistoli et al. 2013). They tend to accumulate in long-lived tissues because of irreversible formation and limited clearance. In diseases such as diabetes and renal failure, the accumulation of AGEs is accelerated and lifestyle factors such as smoking and diet also contribute to the accumulation of a cross-links in extracellular matrix or binding to its transmembrane receptor RAGE. Several studies have found some evidence of an association between AGEs and type 2 diabetes and complications, cardiovascular diseases, and neurodegenerative diseases (Chaudhuri et al. 2013). However, large-scale population based studies are scarce. Within the Rotterdam Study - a large population-based prospective cohort study in the Netherlands - we have assessed AGEs accumulation level in the skin as a reflection of AGEs accumulation in long-lived tissues using a device called the AGE ReaderTM. It measures the skin fluorescence based on the fluorescent property of several AGEs and so far 3009 participants had the measurement from 2013-2016. We have shown cross-sectional subcers and relation with skin AGEs and several traits including vitamin D levels (Chen J et al. 2018), bone fractures (Wagas K 2020), cognition (Chen J et al unpublished, Mooldijk et al 2020) and cardiovascular diseases (Chen J et al. 2020) and several traits including vitamin D levels (Chen J et al. 2018), bone fractures (Wagas K 2020), cognition (Chen J et al. 2020) and with stool microbiome (Chen J et al. 2
Requirements of candidate:	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills. Master degree or MD Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) English language requirement: English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

Dept of Internal Medicine – Genetics Lab & Population Genomics

School/Department:	Department of Internal Medicine-Genetics Lab & Population Genomics, Erasmus MC	
Supervisor information:	Prof. Dr. Joyce B.J. van Meurs (j.vanmeurs@erasmusmc.nl)	
	• Dr. Cindy Boer (c.boer@erasmusmc.nl) Postdoctoral researcher	
world no 29 Endocrinology &	•Website: http://www.glimdna.org ; https://www.linkedin.com/in/joyce-van-meurs-	
<u>Metabolism</u>	78171313/; https://www.erasmusmc.nl/en/research/researchers/meurs-joyce-van	
	•Key words: Population genomics, novel analytic techniques, international and	
	multidisciplinary collaboration, learning environment	
	•Grants:	
	 NWO-VIDI (prestigious Dutch personal grant): €900K) 	
	 H2020 EU: €1500K of in total €12000K 	
	 National Heart, Lung and blood institute (NIH, USA):\$350K of in total \$5000K 	
	- BBMRI-NL roadmap: €2500K	
	 Multiple ZONMW-grants (Dutch Government funding scheme) In total >€1000K 	
	- Erasmus strategic grant: €500K	
	Most important publications:	
	Cell 2021 184:4784-4818 (2021) IF: 38.6] Ann Rheum Dis 2020 80:367-375) [IF:12.4] Ann Rheum Dis 2020 80:598-604) (2021) [IF:12.4] Nat Commun. 2019 Oct 25;10(1):4881. [IF:11.9]	
	Genome Biol. 2019 Nov 14;20:235 [IF:13.2] Nature. 2017 Jan 5;541(7635):81-86. [IF:41.6]	
	Nat Genet. 2017 Jan;49(1):131-138. [IF:27.1] Nat Genet. 2017 Jan;49(1):139-145. [IF:27.1] Nat Commun. 2015;6 [IF14:11.3] Proc Natl Acad Sci, 2012 22;109(21):8218-23 [IF:9.9]	
	Lancet. 2010 Jul 17;376(9736):180-8 [IF: 33.6]	
Project Title:	Large scale population genomics to unravel mechanisms of locomotor diseases	
Abstract:	The Genetic Laboratory of the Department of Internal Medicine has a longstanding tradition	
	and reputation in genomics research, positioned as one of the leading centers in the field of	
	genomics of complex diseases worldwide, with particular focus on locomotor diseases. Prof.	
	Joyce van Meurs has excellent track record in population genetics and genomics studies in	
	osteoarthritis, chronic pain and biological aging. We offer an interesting and challenging	
	position in a multidisciplinary research environment.	
	The project focusses on combining and examining multiple molecular level data	
	((epi)genetics, transcriptomics, proteomics, metabolomics, microbiome) to understand	
	mechanisms of diseases of the locomotor system, such as chronic pain and osteoarthritis.	
	The hallmark of population genomics research is the agnostic, large-scale nature of the data,	
	which allows for novel biological pathways to be discovered. The project is embedded within	
	well-known large scale population studies (Rotterdam Study and Generation R), which have	
	comprehensive phenotyping (including detailed imaging data) as well as a wealth of molecular data available. We also have full access to the UK-biobank data a frequently utilized	
	database for genomics studies. Research will take place in multidisciplinary international	
	consortia, in which the group is well-known and has a leading role. You will explore the	
	available molecular and detailed phenotype data using state-of-the-art analysis techniques	
	(including machine-learning/AI/MR).	
	The aim is to translate the findings of our population genomics studies into two directions:	
	1. Mechanic studies where cell models are used to further study the identified mechanisms;	
	this includes using IPS-cells as a personalized model for disease (done in collaboration with	
	cell biology lab) 2. Application of novel findings into clinic in collaboration with clinical	
	departments.	
Denvironsets of	We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in	
Requirements of	using team work to tackle large scientific questions and thus requires a student with good communication skills.	
candidate:	Master degree or MD	
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) 	
	English language requirement:	
	• English speaking countries & Netherlands: no requirement	
	O Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)	

Dept of Internal Medicine – Genetics Lab & Population Genomics

School/Department:	Department of Internal Medicine-Genetics Lab & Population Genomics, Erasmus MC		
Supervisor information:	Prof. Fernando Rivadeneira (<u>f.rivadeneira@erasmusmc.nl</u>), Full Professor		
	Dr. Ling Oei (<u>h.l.d.w.oei@erasmusmc.nl</u>), Assistant Professor		
erc	Dr. M. Carolina Medina Gomez (<u>m.medinagomez@erasmusmc.nl</u>), Post-doctoral Scholar		
	Website: <u>http://glimdna.org</u> Grants:		
world no 29 Endocrinology &	 Grants: ERC Advanced Grant 2021: €2,500K 		
Metabolism	 Coordinating center European Commission-FP7: HEALTH-2007: €3,000K 		
	- Co-Principal investigator/subcontractor US Government-NIH/R01 2010: \$150K of \$2,500K		
	- Netherlands Consortium of Healthy Aging (NCHA): 2009-2012: €200K		
	- Project manager NWO GROOT Investeringen 2006: €6,000K		
	- NWO VIDI €800K		
	 EU European cooperation in science and technology €150K Marie Skłodowska-Curie Innovative Training Network €520K of €3 800K 		
 Marie Skłodowska-Curie Innovative Training Network €520K of €3,800K Erasmus MC fellowship €400K 			
	Most important publications:		
	2008: Lancet, 371(9623): p. 1505-12. IF:38.3 2009: Nat Genet 41, 1199-206. IF:36.4		
	2010: Nature 467, 832-8 IF:36.3 2012: PLoS Genet, Jul;8(7):e1002718. Epub 2012 Jul 5 IF:9.5		
	2012: Nature Genetics;44(5):491-501. IF:35.2 2012: Diabetes Care;36(6):1619-28. IF:8.57		
	2016: J Bone Miner Res;31(5):1099-106. IF:6.3 2017: Nat Commun;8(1):121. IF: 12.4		
	2018: Am J Hum Genet;102(1):88-102. IF: 9.9 2018: BMJ;362:k3225. IF:27.6 2019: Diabetes Care; 43(1):137-144. IF: 13.4 2018: BMJ;362:k3225. IF:27.6		
Project Title:	Osteoporosis and Environmental Pollution assessed by a Multi-system Approach		
Abstract:			
Abstract:	The Genetic Laboratory of the Department of Internal Medicine has a longstanding tradition		
	and reputation in genomics research and epidemiology, positioned as one of the leading		
	centers in the field of genomics of complex diseases worldwide, with particular focus on		
	musculoskeletal diseases. Our approach is multidisciplinary, combining epidemiology with		
	large-scale genomic and (more recently) microbiome research. The lab is also home to the		
	Generation R and Rotterdam Study cohorts and coordinates the EU-Funded Genetic Factors		
	for Osteoporosis Consortium (GEFOS) consortium and the GEnomics of MusculoSkeletal traits		
	TranslatiOnal expertise Network (GEMSTONE). Prof. Fernando Rivadeneira has excellent track		
	record in genome-wide association studies (GWAS), the epidemiology of diabetic bone		
	disease and Mendelian Randomization (MR) studies. We offer an interesting and challenging		
position in an ambitious yet friendly scientific and clinical research environment			
	(http://glimdna.org).		
	PhD project:		
	You will investigate the influence of environmental pollutants in bone health, through the		
	assessment of endocrine-disrupting chemicals in clinically recruited osteoporosis patients.		
	These individuals will also receive extensive radiological scans and hormone tests in a multi-		
	omic approach, to study the potential underlying pathophysiological mechanisms in different		
	organ systems. Also, questionnaires are collected to potentially advise on healthy lifestyle.		
	Data will be analyzed with both conventional statistics and explorative advanced techniques.		
	Further, collaborative side-projects are possible, including: genetics of diabetic bone disease		
	in type 2 diabetes mellitus in big datasets from population-based studies and clinical cohorts,		
	the potential role of the gut microbiome in the relation of type 2 diabetes and bone disease,		
Requirements of	 clinical risk prediction from polygenic risk scores for various diseases. We are looking for a highly motivated, hardworking student to join our very international team. Our 		
	strength is in using team work to tackle large scientific questions and thus requires a student with good		
candidate:	communication skills.		
	Master degree or MD		
	Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could		
	help with the scientific part of your scholarship proposal)		
	English language requirement:		
	 English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs) TOEEL 100 (min 20 for all subs) 		
	Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)		

Dept of Internal Medicine – Laboratory of Nephrology & Transplantation

The research performed at The Rotterdam Transplantation Laboratory is translational of nature and can be dissected into three research lines being *Transplantation Immunology, Molecular Markers* and *Tissue Repair & Cell Therapy*. Examples of our equipment and operational techniques are: multi-parameter flow cytometry/flow cytometry based cell sorting, imaging flow cytometry, divers cell culture assays i.e., kidney organoids, Elispot, cytotoxicity assays, GWAS, extracellular vesicles, RT-qPCR, epigenetics, histology and immunohistochemistry.





<u>Transplantation Immunology</u>: The wide range of assays to monitor pathways of donor directed reactivity is used to unravel the T and B cell mediated immune responses in patients. In addition, we study the mode of action of (novel) immunosuppressive drugs with the aim to titrate the immunosuppressive burden on our patients in such a way that side-effects (infections, malignancies, cardiovascular events) are kept at a minimum while at the same time rejection processes are prevented.

<u>Molecular Markers</u>: Within this research line we focus on the discovery of molecular markers for either diagnostic or prognostic purposes. We aim to identify patients with complications after kidney transplantation (graft rejection or development of malignancies) in a minimally invasive way via molecular markers in blood or urine. Cell damage due to allograft rejection is accompanied by the release of donor-derived cell-free DNA, extracellular vesicles, and endothelial cells in blood.

<u>Tissue repair & Cell Therapy</u>: We study repair of diseased (transplant) organs by use of cellular therapies such as mesenchymal stem cells. These cells can suppress devastating immune responses against injured organs and stimulate cells within the organs to proliferate and differentiate. Furthermore, we are working on the generation of miniature kidney tissue, so called organoids, from primitive stem cells, which may one day be implanted in the diseased kidney. The aim of these studies is to improve the quality of transplant organs and to repair diseased organs to delay the need for transplantation.

Publications by the Rotterdam Transplant Laboratory

- Shankar AS, et al. Human kidney organoids produce functional renin. Kidney Int 2020 Sep 9:S0085-2538
- Niu Q, et al. Immunosuppression Has Long-Lasting Effects on Circulating Follicular Regulatory T Cells in Kidney Transplant Recipients. Front Immunol. 2020 Aug 28;11:1972.
- Shankar AS, Hoorn EJ, Gribnau J, Baan CC, Hoogduijn MJ. Current State of Renal Regenerative Therapies.Transplantation.2019;103(2):250.
- Yan L, et al. T Follicular Helper Cells As a New Target for Immunosuppressive Therapies. Front Immunol. 2017;8:1510.
- Verhoeven JGHP, et al. Liquid Biopsies to Monitor Solid Organ Transplant Function: A Review of New Biomarkers. Ther Drug Monit. 2018;40(5):515.
- Gonçalves FDC, et al. Membrane particles generated from mesenchymal stromal cells modulate immune responses by selective targeting of pro-inflammatory monocytes. Sci Rep. 2017;7(1):12100

Contact information:

Prof Carla Baan, <u>c.c.baan@erasmusmc.nl</u>, WeChat: carla baan Dr Martin Hoogduijn, <u>m.hoogduijn@erasmusmc.nl</u>, Web: <u>www.RotterdamTransplantationLab.nl</u>

Dept of Internal Medicine – Laboratory of Nephrology & Transplantation

School/Department:	Department of Internal Medicine-Nephrology & Transplantation, Erasmus MC	
Supervisor information:	Prof dr Carla C. Baan (female)	
	Email: c.c.baan@erasmusmc.nl , WeChat:	carla baan
world no 31 Immunology	Website:	Most important publications:.
COLORADO COL	www.rotterdamtransplantationlab.nl	Front Immunol. 2020 Jul 3;11:1332. IF 5.0
电网络苏格里	http://nl.linkedin.com/pub/carla-baan/8/a19/960	Front Immunol. 2020 Aug 28;11:1972. IF 5.0
64 - A - A - A - A - A - A - A - A - A -	www.erasmusmc.nl	Drugs. 2020 Jan;80(1):33-46. IF 6.2
(199) (1992	Personal Grants:	Kidney Int 2020Sep 9:S0085-2538(20)30968-6. IF 8.4
2.22394230	2019, Dutch Kidney Foundation	Transplantation. 2020 Mar 6. IF 4.5
前的影响中的	2018, Astallas Pharma 2017, Dutch Kidney Foundation	<u>Transplantation 2019 May;103(5):e110-e111</u> . IF 4.5 <u>Ther Drug Monit. 2018;40(5):515-525</u> . IF 2.4
	2016, Lundbeck Foundation Denmark	Front Immunol. 2017;8:306. IF 6.5
		<u>Sci Rep. 2017;7:12100</u> . IF 4.1
Project Title:	le: Exploiting the message from the kidney: the value of extracellular vesicles	
	transplant rejection	
Abstract:	Worldwide, approximately 80.000 kidney tran	splantations are performed annually. Without a
		d, and immune competent cells like T cells will
		o 25% of cases, but the reasons for rejection are
	still largely unknown. The discovery that extra	· · · · · · · · · · · · · · · · · · ·
		s and that they readily cross cell walls is a boon
		and cellular mechanisms driving this response.
	We propose the novel concept that donor or	0
	way for recipient immune cells to initiate the	
	novel ex vivo platform will be developed to de	
	delivery of extracellular vesicle cargo to immu	
		man cells and transport cell-derived molecules
	to other cells, changing their phenotype and f	•
	extracellular vesicles	
	H-MHC	Secreting Cell
	· · · ·	15
	I UTEIgH alltigens	Blebbing Apoptotic Bodies
	including the immune	
	activating proteins that	Early Endosome
	interact with recipient	MVB
	antigen presenting cells Shedding	Budding J Exocytosis
	and sets off the T cell	
	dominated immune Microvesicles	Cytosol
	response. Technological	Exosomes
	advances in ex vivo	· · · · · · · ·
	tissue engineering	
	systems, imaging	
	technologies and omics	
	now facilitate the study	Recipient Cells
	of 1) how donor kidney-extracellular vesicles interact with recipient antigen presenting cells,	
		at means we can interfere in this reaction. This
	study delivers new knowledge about immune	-
	importance in auto-immunity, cancer and infe	
Requirements of		student to join our very international team. Our strength is in and thus requires a student with good communication skills.
candidate:	Master degree or MD	and this requires a student with good communication skills.
	Scholarship that will, at least, cover subsistence allo	wance and international air plane ticket (we could help with the
	scientific part of your scholarship proposal)	
	 English language requirement: English speaking countries & Netherlands: no requir 	ement
	Lingion openning countries & netherlands. no requir	FL 100 (min 20 for all subs)

Dept of Internal Medicine – Laboratory of Nephrology & Transplantation

School/Department:	Department of Internal Medicine-Nephrology & Transplantation Erasmus MC		
Supervisor information:	Dr Martin J Hoogduijn		
	Email: <u>m.hoogduijn@erasmusmc.nl</u>		
world no 31 Immunology	Website:		
	- https://loop.frontiersin.org/people/29382/overview		
	- <u>https://www.rotterdamtransplantationlab.nl/</u>		
	Personal Grants:		
	 2018 - 2021 Health Holland TKI grant 2016 - 2020 Lundbeck Foundation Denmark 		
	- 2014 - 2018 FP7 EC project		
	Most important publications:		
	- Shankar et al. Transplantation 2019 Vol 103(2);250-261		
	- De Witte et al. Stem Cells. 2018 Vol 36(4);602-615		
	 Goncalves et al. Scientific Reports 2017 Vol 21;7(1):12100 Hoogduijn et al. British Med J 2013 Vol 347:f6833 		
	 Eggenhofer et al. Front Immunol. 2012 Vol 3:297 		
Project Title:	Generation of kidney organoids from pluripotent stem cells		
-			
Abstract:	Regenerative medicine holds potential to cure multiple diseases, including kidney disease. In		
	recent years protocols were developed to differentiate human induced pluripotent stem cells (iPSC) into kidney organoids. Using these protocols, kidney organoids can be generated with		
	distinct tubular and glomerular structures that can survive for several months after in vivo		
	implantation (Figure below). However, the kidney organoids generated with the current		
	protocols are immature and resemble kidneys of first trimester embryos. Further maturation		
	of kidney organoids is needed to make them suitable for kidney disease modeling, drug testing		
	and eventually for repair of lost kidney function.		
	In this project we will improve the maturation status of kidney organoids by introducing		
	adaptations in the culture protocol and subsequent implantations of the organoids under the		
	skin or into the injured kidney. The maturation status of kidney organoids will be examined		
	morphologically by (confocal) microscopy, through gene and protein expression analysis using		
single cell sequencing and proteomics. The functionality of the organoids will be det by examining their urine production function and their capacity to produce hormone			
			secreted by the adult kidney.
	Example of immunohistochemically stained kidney organoid		
	generated from human iPSC in vitro (left) and zoomed in		
	glomerulus of a kidney organoid subcutaneously implanted		
Requirements of	 in a mouse (right). We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using 		
candidate:	team work to tackle large scientific questions and thus requires a student with good communication skills.		
cunululle.	 Master degree or MD Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we will help with the 		
	scientific part of your scholarship proposal)		
	English language requirement: - English speaking countries & Netherlands: no requirement		
	- English speaking countries & Netherlands: no requirement - Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)		

School/Department:	Department of Internal Medicine-Metabolism & Reproduction, Erasmus MC
Supervisor information:	• Dr. Ir. Jenny A. Visser
	• Email: j.visser@erasmusmc.nl
world no 29 Endocrinology &	Website: <u>https://www.erasmusmc.nl/en/research/groups/metabolism-and-reproduction</u>
Metabolism	https://www.linkedin.com/in/jenny-visser-1375357/
	• Grants:
	 2019 - 2022 Health Holland TKI grant Royalties
	Most important publications:
	 Hoyos LR et al. Loss of anti-Müllerian hormone (AMH) immunoactivity due to a homozygous AMH gene variant rs10417628 in a woman with classical polycystic ovary syndrome (PCOS). Hum Reprod. 2020, 35(10):2294-2302. Moolhuijsen LME, Visser JA. Anti-Müllerian Hormone and Ovarian Reserve: Update on Assessing Ovarian Function. J Clin
	 Endocrinol Metab. 2020, 105(11):dgaa513. Kaikaew K et al. Sex Difference in Corticosterone-Induced Insulin Resistance in Mice. Endocrinology. 2019, 160(10):2367-
	 2387. Day F et al. Large-scale genome-wide meta-analysis of polycystic ovary syndrome suggests shared genetic architecture for different diagnosis criteria. PLoS Genet. 2018, 14(12):e1007813.
	- Day FR et al. Genomic analyses identify hundreds of variants associated with age at menarche and support a role for puberty timing in cancer risk. Nat Genet. 2017, 49(6):834-841.
	 Mahfouz A et al. Genome-wide coexpression of steroid receptors in the mouse brain: Identifying signaling pathways and functionally coordinated regions. Proc Natl Acad Sci U S A. 2016, 113(10):2738-43.
	 Day FR et al. Large-scale genomic analyses link reproductive aging to hypothalamic signaling, breast cancer susceptibility and BRCA1-mediated DNA repair. Nat Genet. 2015, 47(11):1294-1303. Grefhorst A et al. Estrogens increase expression of bone morphogenetic protein 8b in brown adipose tissue of mice. Biol Sex
	Differ. 2015,6:7.
	- van Houten E et al.Reproductive and metabolic phenotype of a mouse model of PCOS. Endocrinology. 2012, 153(6):2861-9.
Project Title:	Understanding sex differences in metabolism
Abstract:	Obesity remains a prevalent global public health issue as it is a major risk factor for type 2
	diabetes, cardiovascular diseases and cancer. Although the global prevalence of obesity is
	higher in women than in men, obese men are more prone to develop obesity-related
	conditions than obese women. This sex difference diminishes when women enter
	menopause, suggesting a prominent role for sex steroids in controlling metabolism. Indeed,
	disturbances in gonadal function are associated with metabolic problems. For instance,
	obesity and insulin resistance is frequently present in women with polycystic ovary syndrome (PCOS), a disease characterized by hyperandrogenism.
	Our studies are aimed at understanding the mechanisms that contribute of the sexual
	dimorphism in metabolic diseases. We have several research projects in which we delineate
	the effects of altered sex steroids and gonadal growth factors (such as AMH) on metabolism.
	In particular, we aim to understand why the effects of sex steroid hormones differ in male vs
	female white and brown adipose tissues. We also study how gut hormones contribute to sex
	differences in metabolism. Studies are performed at physiological (mouse models), cellular
	(iPS cells), and molecular level. In addition, studies will be performed at a genetic level in
	collaboration with (inter)national consortia.
Requirements of candidate:	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills. Master degree or MD (with experience in molecular biology techniques) Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) Explicit largue requirement.
	English language requirement: <i>English speaking countries & Netherlands:</i> no requirement
	 Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

Dept of Internal Medicine – Neuroendocrine Tumors

School/Department	Dept Internal Medicine - Neuroendocrine Tumors, Erasmus MC
Supervisor	Prof. Dr. W.W. de Herder & Dr. J. Hofland
information:	• Email: w.w.deherder@erasmusmc.nl & j.hofland@erasmusmc.nl
	Website: <u>https://www.erasmusmc.nl/en/research/departments/internal-medicine-laboratories</u>
world no 20	Personal Grants:
world no 29	• ERC H2020 Marie-Curie Intra-European Fellowship (2013), Royal College of Physicians UK (2013), Daniel den Hoed Foundation
Endocrinology &	(2015), Erasmus MC MRACE-Grant (2017), Swiss National Science Foundation (2018), co-investigator Dutch Cancer Fund (2019),
<u>Metabolism</u>	NET Research Foundation (2020)
	 Most important publications: Additional holmium-166 radioembolisation after lutetium-177-dotatate in patients with neuroendocrine tumour liver metastases
	(HEPAR PLuS): a single-centre, single-arm, open-label, phase 2 study. Lancet Oncol 2020; 21: 561-570
	 Advances in the diagnosis and management of well-differentiated neuroendocrine neoplasms. Endocr Rev 2020; 41: 371-403
	• Management of carcinoid syndrome: a systematic review and meta-analysis. Endocr Relat Cancer. 2019; 26: R145-156
	• Symptomatic and radiological response to 177Lu-DOTATATE for the treatment of functioning pancreatic neuroendocrine tumors. J
	Clin Endocrinol Metab 2019, 104(4): 1336-1344
	Salvage peptide receptor radionuclide therapy with [177Lu-DOTA,Tyr3]octreotate in patients with bronchial and astroantereparateratic pourceptore tumours. Fuel Nucl Med Med Imaging 2010, 46(2):704,717
	 gastroenteropancreatic neuroendocrine tumours. Eur J Nucl Med Mol Imaging 2019, 46(3):704-717. Role of biomarker tests for diagnosis of neuroendocrine tumours. Nature Rev Endo 2018, 14(11):656-669
	 MAFA missense mutation causes familial insulinomatosis and diabetes mellitus. PNAS 2018 Jan 30;115(5):1027-1032
	Persistent Hematologic Dysfunction after Peptide Receptor Radionuclide Therapy with 177Lu-DOTATATE: Incidence, Course, and
	Predicting Factors in Patients with Gastroenteropancreatic Neuroendocrine Tumors. J Nucl Med. 2018 Mar;59(3):452-458
	Consensus on biomarkers for neuroendocrine tumour disease. Lancet Oncol. 2015 Sep;16(9):e435-e446.
Project Title:	Discovery of novel biomarkers for gastroenteropancreatic neuroendocrine tumors
Abstract:	Neuroendocrine neoplasms of the pulmonary and gastrointestinal systems are heterogeneous
	tumors. Although rare, their incidence has risen 6-fold over the last 3 decades. Well-differentiated
	neuroendocrine tumors (NETs) have limited treatment options and are often accompanied by severe
	hormonal syndromes. Our NET Center of Excellence has been world-leading in this field with
	translational biomarker research ^(Nature Rev Endo 2018) , participation in international
	guidelines ^(Neuroendocrinology 2016) and the development of radionuclide imaging ^(Lancet 1989) and therapy ^{(NEJM}
	2017)
	Our research lines in endocrine oncology have a strong translational aspect with close interaction
	between clinical and basic scientists. We participate in international clinical trials, have created
	clinical databases with >2000 NET patients and have a dedicated Neuroendocrine Laboratory with
	decades of experience in in vitro and ex vivo characterization of NET cells.
	Current projects focus on the discovery of novel biomarkers for gastroenteropancreatic NETs
	through epigenomics, proteomics and microbiomics. This includes regulatory control of
	somatostatin receptor expression as well as the search for biomarkers for carcinoid syndrome-
	related complications and for the efficacy of peptide receptor radionuclide therapy (PRRT). This
	project will integrate into our long-standing translational biomarkers studies to improve diagnostics,
	prognostication and prediction of therapeutic outcome in patients with bronchial and
	gastroenteropancreatic NETs.
Requirements of	• We are looking for a highly motivated and enthusiastic student to join our international team. The candidate should be a team
candidate:	player with good communication and writing skills and interested in translational cancer science
	 Master degree or Medical Degree. Prior experience in molecular biology, bioinformatics and statistics is of significant added value. Scholarchin that will at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part).
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal)
	 English language requirement: fluently speaking and writing.
	English speaking countries & Netherlands: no requirement
	• Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

Dept of Internal Medicine – Thyroid Function in Health & Disease

School/Department:	Department of Internal Medicine-Thyroid Function in Health & Disease, Erasmus MC
Supervisor information:	Prof dr R.P. Peeters & Dr. W.E. Visser
	Email: <u>r.peeters@erasmusmc.nl</u> & <u>w.e.visser@erasmusmc.nl</u>
world no 29 Endocrinology &	Website: <u>https://www6.erasmusmc.nl/inwendige_geneeskunde/endocrinologie/research</u>
<u>Metabolism</u>	Personal Grants:
	- ZonMW VENI grant and VIDI grant (Dutch equivalents of ERC Starting and Advanced Grant),
	- ZonMW Clinical Fellowship, - ZonMW TOP Grant,
	- and several EU-Horizon2020 Grants
	Most important publications:
	 Peeters RP. Subclinical Hypothyroidism. N Engl J Med. 2017 376(26):2556-2565 & N Engl J Med. 2017 377(14):1404. Korevaar TIM, Medici M, Visser TJ, Peeters RP. Thyroid disease in pregnancy: new insights in diagnosis and clinical management. Nature Rev Endocrinol. 2017 13(10):610-622.
	- Chaker L, Bianco AC, Jonklaas J, Peeters RP. Hypothyroidism. Lancet. 2017
	 Teumer A, Chaker L, Groeneweg S,, Peeters RP, Naitza S, Völzke H, Sanna S, Köttgen A, Visser TJ, Medici M. Genome-wide analyses identify a role for SLC17A4 and AADAT in thyroid hormone regulation. Nature Commun. 2018 Oct 26;9(1):4455. Maternal thyroid function during pregnancy and child brain morphology: a time window-specific analysis of a prospective of the second s
	cohort. Jansen TA, Korevaar TIM, Mulder TA, White T, Muetzel RL, Peeters RP, Tiemeier H. Lancet Diabetes Endocrinol. 2019 Aug;7(8):629-637.390(10101):1550-1562.
	 Effectiveness and safety of the tri-iodothyronine analogue Triac in children and adults with MCT8 deficiency: an international, single-arm, open-label, phase 2 trial. Groeneweg S, Peeters RP, Moran C,, Polak M, Chatterjee K, Visser TJ, Visser WE. Lancet Diabetes Endocrinol. 2019 Sep;7(9):695-706.58
	- Association of Thyroid Function Test Abnormalities and Thyroid Autoimmunity With Preterm Birth: A Systematic Review and
	Meta-analysis. Consortium on Thyroid and Pregnancy—Study Group on Preterm Birth, Korevaar TIM, Derakhshan A, Taylor PN,
	Meima M,, Steegers EAP, Peeters RP. JAMA. 2019 Aug 20;322(7):632-641
Project Title:	Consequences of thyroid dysfunction for development, metabolism and aging
Abstract:	Thyroid hormone is essential for normal growth, metabolism and adequate functioning of
	almost all tissue. Thyroid dysfunction is a very prevalent disorder, with hypothyroidism
	affecting circa 5% of the population. It is more prevalent in women and in elderly.
	We study the consequences of disturbances of thyroid hormone action at multiple levels. In
	close collaboration with the department of epidemiology, we study the consequences of mild
	alterations in thyroid function on child development (Lancet Diab and Endo 2019) and
	pregnancy outcome (JAMA 2019) in the large population-based birth cohort Generation R,
	whereas we study the consequences of thyroid dysfunction on the aging process (JAMA
	Intern Med 2017 & Circ Res 2017) in the population-based Rotterdam Study. We closely
	collaborate with other renowned population-based studies across Europe and United States
	and initiated two consortia (JAMA 2019 & Nature Communications 2018).
	In addition, we have several research projects in which we delineate the consequences of
	genetic defects in thyroid hormone pathways genes at the molecular level. This led to the
	identification of different types of thyroid hormone insensitivity due to defects at the level of
	uptake into the cell (MCT8 deficiency, Lancet 2004) or at the receptor level (NEJM 2012). The
	studies performed in this area focus on understanding the molecular mechanisms leading to
	these diseases, as well as developing treatments. This has led to the first international clinical
	trial for MCT8 deficiency (Lancet Diab & Endo), which was coordinated by our group.
Requirements of	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to taskle long scientific questions and thus requires a student with good communication skills.
candidate:	 using team work to tackle large scientific questions and thus requires a student with good communication skills. Master degree or MD (with experience in molecular biology techniques)
	• Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the
	scientific part of your scholarship proposal)
	English language requirement: Fraglish spaceting countries & Netherlands, no requirement
	 English speaking countries & Netherlands: no requirement

Department of Medical Oncology

The treatment of an individual with cancer is determined by specific characteristics of that individual patient, the cancer cells, and their environment, and needs to be constantly adjusted according to the changes observed in these characteristics. To improve treatment, we need to improve our understanding of the many characteristics determining the outcome of patients after treatment. Three of our key research areas are:

Translational Cancer Genomics and Proteomics (PI Prof. Dr. John Martens)

We aim to discover clinically relevant breast, colorectal and prostate cancer biomarkers of disease progression using genomics techniques.

- We use various genomics tools (RNA sequencing; next generation sequencing) to discover and validate new prognostic and predictive markers providing insight into molecular mechanisms of disease progression and therapy failure. It is our ambition to offer patients the best possible choice of treatment.
- To understand the evolution of metastatic cancer towards therapy resistance we study the temporal variation in various types of circulating biomarkers (circulating tumor cells (CTCs) and circulating endothelial cells (CECs); circulating nucleic acids (ctDNA/ctRNA) and exosomes) during therapy.

Key publications

- 1. Smid M et al. Breast cancer genome and transcriptome integration implicates specific mutational signatures with immune cell infiltration. Nat Commun. 2016; 7:12910.
- 2. Sieuwerts AM, et al. mRNA and microRNA expression profiles in circulating tumor cells and primary tumors of metastatic breast cancer patients. Clin Cancer Res. 2011 17:3600-3618.
- 3. Angus L, et al. Genomic landscape of a large cohort of metastatic breast cancer patients. Nat. Genetics. 2019.

Translational Immuno-Oncology (PI Assoc Prof Dr. Reno Debets)

We aim to understand T cell immunity in common tumor types and enable treatment of patients with customized combination adoptive T cell therapy. To this end, we follow 3 research lines:

- Develop and test adoptive T cell therapy: selection and validation of targets and receptors, gene-engineering of T cells, and implementation of clinical T cell treatments (>15-year track record). Our laboratory has tested gene-engineered T cells in advanced renal cell cancer, the 1st clinical study of its nature in Europe (completed). We are currently selecting safe and effective targets and obtaining corresponding TCRs according to a stepwise approach using the latest in silico and laboratory tools: a first product (a TCR against MAGE-C2) is scheduled for clinical testing in Q4 2019.
- Understand and intervene with T cell immunity: discovery and functional assessment of determinants of anti-tumor T cell immunity using techniques that address frequencies, functions and spatio-organization of T cells as well as intervention studies with (immune) modulators using 3D cultures and syngeneic and immune deficient mouse models.
- Monitor patient T cell immunity: we phenotypically assess changes of T cell (subsets) in blood and tissue of patients with various tumor types in relation to resistance to (immune)therapies, to stratify patients and guide selections of drugs that make tumors better amenable to T cell treatments.

Key publications

- 1. Straetemans T et al. Recurrence of melanoma following T cell treatment: continued antigen expression in a tumor that evades T cell recruitment. Mol Ther. 2015 23:396.
- 2. Hammerl D et al. Adoptive T Cell Therapy: New Avenues Leading to Safe Targets and Powerful Allies. Trends Immunol, 2018 18:30169.
- 3. Kunert A et al. CD45RA+CCR7- CD8 T cells lacking co-stimulatory receptors demonstrate enhanced frequency in NSCLC patients responding to nivolumab. J Immunotherapy Cancer, 2019 7:149.

Prostate Cancer Clinical Trials (PI Dr. Martijn Lolkema)

- Genomic classification of prostate cancer patients to predict outcome to anti-cancer treatment. In collaboration with the Hartwig
 Medical Foundation and the Center for Personalized Cancer Treatment we obtained Whole Genome Sequencing data from > 400
 prostate cancer patients and we are analyzing the data in order to understand the inter-patient heterogeneity. Moreover, we are
 building a biobank of clinically annotated samples (circulating markers and tissue biopsies) from patients with metastatic prostate
 cancer who are actively undergoing treatment.
- Prospective Clinical Trials. We perform prospective clinical trials in prostate cancer patients mainly based on biomarker stratification such as a trial in which we use patient selection using AR-V7 expression in CTCs to allocate patients for cabazitaxel treatment.

Key publications

- 1. Van Dessel et al. The genomic landscape of metastatic castration-resistant prostate cancers using whole genome sequencing reveals multiple distinct genotypes with potential clinical impact https://www.biorxiv.org/content/10.1101/546051v1
- 2. Belderbos et al. Associations between AR-V7 status in circulating tumour cells, circulating tumour cell count and survival in men with metastatic castration-resistant prostate cancer. Eur J Cancer. 2019 121:48-54.
- 3. Priestley et al. Pan-cancer whole genome analyses of metastatic solid tumors. https://www.biorxiv.org/content/10.1101/415133v4

Department of Medical Oncology

School/Department:	Department of Medical Oncology Erasmus MC
Supervisor information: world no 32 Oncology	 Prof dr. John Martens (supervisor) Dr. Harmen van de Werken (co-supervisor) Email: j.martens@erasmusmc.nl and/or h.vandewerken@erasmusmc.nl Website: John Martens and Harmen van de Werken & [] Personal Grants: DDHF CCBC (2014 & 2018) Astellas (ML; 2014) NKB EMCR (2014) Most important recent publications: Lindsay Angus,, Harmen J.G. van de Werken ,, John W.M. Martens 2019. "Genomic landscape of metastatic breast cancer and its clinical implications". Nature Genetics 51(10):1450-1452. Harmen J.G. van de Werken, van Riet, J.*, and Mostert, B. 2021 The genomic landscape of Sa dvanced neuroendocrine neoplasms reveals subtype-heterogeneity and potential therapeutic targets. Nature Communications 12, 1–14. Nik-Zainal, Serena,, John W. M. Martens,, and Michael R. Stratton. 2016. "Landscape of Somatic Mutations in 560 Breast Cancer Whole-Genome Sequences." Mature 3347605;147–54. Smid, Marcel,, John W. M. Martens,, and Michael R. Stratton. 2016. "Landscape of Somatic Mutations in 560 Breast Cancer Whole-Genome Sequences." Mature Communications 7:12910. Harmen J.G. van de Werken et al 2017 Small chromosomal regions position themselves autonomously according to their chromatin class. Genome Res. 27, 922–933 van de Werken, Harmen J. G., 2012 et al. "Robust 4C-Seq Data Analysis to Screen for Regulatory DNA Interactions." Mature Methods 9(10):969–72.
Project Title:	Cancer Computational Biology to Gain Insights in Biology and Create Clinical Value Using Multi-Omics Data Sets of Advanced and Metastatic Patients
Abstract:	A Dutch initiative involved the biobanking of tumor biopsies and matched blood samples from cancer patients with locally advanced and metastatic diseases and subjecting them to Whole Genome Sequencing (WGS). The heroic effort generated a database of currently more than 4000 WGS datasets revealing pan-cancer and subtype specific driver events and mutational programs relevant for disease progression and therapy failure. In these first studies matched transcriptomics, in addition to WGS data, were not included as these data were generated at a later time point. Therefore, the next intruding step is to interrogate available transcriptome data and integrate them with matched WGS data. This provides us with the opportunity, in metastatic cancer, 1) to identify the phenotypic heterogeneity, 2) the clinical significance of RNA-seq beyond WGS data 3) and identify novel disease progression and cancer drug-resistances modules. Currently, we have access to 2072 matched RNA-seq datasets from 36 cancer types and eight different treatment categories, including chemotherapy and immunotherapy. We will interrogate this very comprehensive data set by applying state-of-the art- bioinformatic and computational biology methods including regularized multivariate analyses and machine learning methods, such as Random Forest and Neural Networks. The insights we will gain from this interrogation will be incorporated in patient stratification statistical models to ultimately support physicians in their clinical decision making, which may improve the health of cancer patients in the future.
Requirements of candidate:	 We are looking for a candidate with strong analytical and problem-solving skills, being highly motivated and having excellent communication and writing skills and able to work independently. A background in cancer biology is of significant added value. Master's degree in bioinformatics, computational biology, statistics, or a related field. The candidate should have demonstrated excellent scientific writing and software engineering skills in R and Python or Perl. Scholarship that will, at least, cover subsistence allowance and international airplane ticket (we could help with the scientific part of your scholarship proposal) English language requirement: <i>English speaking countries & Netherlands:</i> no requirement <i>Other countries:</i> IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

Department of Medical Oncology

School/Department:	Laboratory of Tumor Immunology, Department of Medical Oncology. Erasmus MC
Supervisor information:	Supervisors:
	Dr. Hayri Emrah Balcioglu (<u>h.balcioglu@erasmusmc.nl</u>)
world no 32 in Oncology	Prof. Dr. Reno Debets (<u>j.debets@erasmusmc.nl</u>)
	Website:
	https://www.erasmusmc.nl/en/cancer-institute/research/groups/medical-oncology-tumor-
	immunology ; https://www.tme-facility.com
	5 grants (out of 15 running grants):
	- Dutch Cancer Society; Adoptive therapy with T cells gene-engineered with a co-stimulatory TCR to treat
	 patients with MAGE-C2-positive melanoma and head and neck cancer. 570 k€. Merck; Genomic and immune profiling of metastasized urothelial cancers.735 k€.
	 Dutch Cancer Society; Co-stimulatory TCRs to advance treatment efficacy of adoptively transferred T cells.
	457 k€.
	 Erasmus MC Daniel den Hoed Foundation; Adoptive T cell therapy to treat common cancers: new roads to unique targets and pre-treatments. 500 k€.
	 Top consortia for knowledge and innovation (Dutch government); T-cells act against hard-to-treat cancers (T-ACT): unique targets and new technological platform to develop safe and effective adoptive cellular therapeutics (T-ACT). 900 k€.
	5 publications (out of 150):
	- Lamers C et al. Treatment of metastatic renal cell carcinoma with autologous T-lymphocytes genetically retargeted against carbonic anhydrase IX: first clinical experience. J Clin Oncol, 2006 24:e20.
	- Straetemans T et al. Recurrence of melanoma following T cell treatment: continued antigen expression in a tumor that evades T cell recruitment. Mol Ther , 2015 23:396.
	- Kunert A et al. <u>T cell receptors for clinical therapy: <i>in vitro</i> assessment of toxicity risk. Clin Cancer Res, 2017 23:6012.</u>
	- Kortleve D et al. News and views: Orthoptopic editing of T-cell receptors. Nature Biomedical Engineering , 2019, 3:949.
	- Hammerl D et al. Spatial immunophenotypes predict resistance to anti-PD1 treatment and capture distinct paths of T-cell evasion in triple negative breast cancer. Nature Comm , in press.
Project Title:	CD8 T-cell trafficking and activity captured in patient 3D spheroid model
Abstract:	Emergence of immunotherapy has changed the treatment and patient outcome for various
	tumor types. Unfortunately, patient response and reasons behind failure of response is
	currently hard to assess. In the laboratory of tumor immunology, we aim to define and
	understand shortcomings of T cell immunity in cancers, and translate our findings into the
	development of anti-cancer T cell treatments. The T cell migration towards tumors, and
	accumulation and activation in the tumor is crucial for the success of immunotherapy. Along
	this line, it is imperative to capture the real dynamics of patient T cell activity, particularly the
	interactions between T cells and tumor cells, or lack there-of.
	Recently, we have set up a 3-D tumoroid model to monitor movement and anti-tumor activity
	of patient T cells in real-time. This technique enables quantification of patient T cell migration,
	infiltration, activation and tumor clearance in 3D. With this project, the PhD candidate will
	determine differences in such dynamics between T cells derived from patient tumors that are
	responsive versus those that are not responsive to immune therapies. In more detail, the
	candidate will study tumor cell-directed mechanisms of T cell suppression, and will correct
	such T cell suppression via genetic and pharmacological means, ultimately, identifying
	determinants of response to therapy, and targets for sensitization of non-responsive tumors
	to immunotherapy.
Requirements of	 highly motivated, hardworking background in cancer biology, mechanobiology and/or tumor immunology is a preferred value
candidate:	- master degree or MD.
	 scholarship that will cover subsistence allowance and international air plane ticket english language requirement:
	• English speaking countries & Netherlands: no requirement
	• Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Molecular Genetics, Erasmus MC
Supervisor information:	Dr. Miao-Ping Chien, m.p.chien@erasmusmc.nl, http://www.mpchienlab.org/
	Selected Grants:
World no 30 Biomedical	2021 Oncode Technology Development Grant 2018 CancerGenomiCs.nl Junior Pl's Grant
<u>Sciences</u>	2020 Ammodo Science Award 2018 Dragon Gate Grant (Taiwan MoST)
	2020 Erasmus-TU Delft Convergence Grant 2017 NWO Veni award (NWO Talent Scheme)
	2019 Oncode Institute Junior Fellow2017 CancerGenomiCs.nl Junior Fellow2018 Erasmus MC Fellowship
	Selected publications:
	1. You, Li*, Su, P.R.*, Betjes, M.*, Ghadiri Rad, R., Chou, T.C., Beerens, C., van Oosten, E., Leufkens, F.,
a corces	Gasecka, P., Muraro، M., van Tol، R., van Steenderen, D., Farooq, S., Hardillo, J.A.O., Baatenburg de
	Jong, R., Brinks, D.A, Chien, M.P. "Functionally annotated transcriptomic profiling of single cells from
	heterogeneous populations based on dynamic phenotypes", Nature Biomedical Engineering , In press
	(2021) 2. Su, P.R., You, L., Beerens, C., Bezstarosti, K., Demmers, J., Pabst, M., Kanaar, R., Hsu, C.C., Chien,
	<i>M.P.</i> , "Functional single cell proteomic profiling of cells with abnormal DNA damage response
Miao-Ping Chien received her	dynamics". Under review
PhD in chemistry and biochemistry from the	3. Li L et al. " <u>A Comprehensive enhancer screen identifies TRAM2 as a key and novel mediator of YAP</u>
biochemistry from the University of California, San	oncogenesis." Genome Biology, 2021, 22, 54,
Diego in 2013, and went on to	4. Chien M.P et al. "Photoactivated voltage imaging in tissue with an archaerhodopsin-derived
do a postdoc at Harvard	 <u>reporter</u>, Science Advances, 2021: Vol. 7, no. 19, eabe3216 Werley C.A., et al <u>"An ultrawidefield microscope for high-speed fluorescence imaging and targeted</u>
University, working on	optogenetic stimulation." Biomedical Optics Express. 2017, 8(12), 5794-5813.
technology development for	6. Chien M.P. , et al. <u>"Enzyme-Directed Assembly of Nanoparticles in Tumors Monitored by In Vivo</u>
biology (combining biophysics,	Whole Animal and Ex Vivo Super Resolution Fluorescence Imaging." J Am Chem Soc. 2013 Dec
computation and optical	18;135(50):18710-3.
instrumentation). She joined	7. Chien M.P., et al. <u>"Enzyme-Directed Assembly of a Nanoparticle Probe in Tumor Tissue."</u> Advanced
Erasmus MC as a group leader	Materials. 2013, July 12 (25): 3599-3604.
in June 2017 and became a	Investigation of tumorigenesis via advanced imaging and single cell -omics
principal investigator at Oncode	analysis
Institute in 2019. Her current	The Chien Lab is looking for self-motivated PhD students with a strong interest in
research focuses on developing	working in a multidisciplinary lab. In our lab, we develop single cell technologies
and applying multidisciplinary	combining optical, biomedical and bioinformatics methods to address biological
technologies (advanced	questions, particularly in cancer biology and immuno-oncology.
microscopy and imaging,	
computation, single cell technology, bioinformatics,	The candidate will have a chance to work on wet-lab projects, dry-lab projects or a
(photo)chemistry) to investigate	combination of these two. For the wet-lab projects, the candidate can apply the
the underlying mechanisms of	technologies developed in Dr. Chien's group, including advanced imaging and single
tumorigenesis, particularly of	cell sequencing (analysis), to cancer cell lines or patient-derived primary cultures to
rare cancer-driving cells. She is	investigate molecular mechanisms of tumorigenesis and therapy resistance. For the
also a founder of UFO	dry-lab projects, the candidate can work on advanced imaging analysis including
Biosciences, which aims to	machine learning-based approaches or bioinformatics analysis (-omics data
enable better cancer care by	
creating treatment options for	analysis).
rare, cancer-driving cell	
populations that escape	
traditional treatment.	
Requirements of candidate:	• We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in
	 using team work to tackle large scientific questions and thus requires a student with good communication skills. Master degree or MD
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the
	scientific part of your scholarship proposal)
	English language requirement: English spagking countries & Netherlands: no requirement
	 English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Molecular Genetics, Erasmus MC
Supervisor information:	• Dr. Hannes Lans, Associate professor DNA repair mechanisms and disease
	• w.lans@erasmusmc.nl www.lanslab.eu
World no 30 Biomedical	
<u>Sciences</u>	• Grants:
	 2018 2x Dutch Research Council (€ 568000)
	- 2017 Dutch Cancer Society (€ 534000)
	 - 2014 WorldWide Cancer Research (€ 218000)
	- 2012 MSCA FP7-PEOPLE-ITN (€ 689000)
	- 2008 Veni grant Dutch Research Council (€ 208000).
	Most important publications:
	Ribeiro-Silva C et al (2020) Ubiquitin and TFIIH-stimulated DDB2 dissociation drives DNA damage
	handover in nucleotide excision repair. <i>Nature Communications</i> 11:4868
	Lans H et al (2019) The DNA damage response to transcription stress. Nature Reviews Mol Cell Biol
	20:766-784
	Borgermann N et al (2019) SUMOvation promotes protective responses to DNA-protein crosslinks.
	EMBO Journal 38:e101496
	Ribeiro-Silva C et al (2018) DNA damage sensitivity of SWI/SNF-deficient cells depends on TFIIH subunit
	p62/GTF2H1. Nature Communications 9:4067
	Slyskova J et al (2018) Base and nucleotide excision repair facilitate resolution of platinum drugs-
	induced transcription blockage. Nucleic Acids Research 46:9537-9549
	Marteijn JA et al (2014) <u>Understanding nucleotide excision repair and its roles in cancer and ageing</u>
	Nature Reviews Mol Cell Biol 15:465-81
Project Title:	Cell-type specific functional analysis of DNA repair
Abstract:	Accumulation of DNA damage is an important underlying cause of major health issues like
	cancer and aging. Nucleotide excision repair (NER) is a major cellular defense mechanism that
	repairs a large variety of helix-distorting DNA damage, including that induced by solar UV
	irradiation and platinum-based anticancer drugs. Hereditary defects in NER cause multiple
	different cancer-prone and degenerative diseases in which tissues are differently affected,
	but of which the exact pathogenesis is not understood. We have found that NER activity
	changes depending on development and cell type, but how this is regulated is not known.
	We investigate the tissue-specific activity of NER through the identification and functional
	characterization of novel regulatory proteins and mechanisms within this important DNA
	repair pathway. To this end, we use different model systems, including <i>C. elegans</i> ,
	mammalian cell culture and <i>in vitro</i> differentiated cells (based on induced pluripotent stem
	cells). We pursue a multi-disciplinary approach, using cell biology, CRISPR- and RNAi-mediated
	screening combined with live cell confocal microscopy and quantitative proteomics, to study
	NER mechanisms in different cell types. We are looking for a highly motivated PhD student
	who wants to work on this frontline ambitious project aimed at understanding how NER
	protects different cell types against DNA damage. The results of this project will help to better
	understand the molecular pathogenesis associated with inherited NER deficiency and to
	develop therapies aimed at alleviating discomfort associated with cancer and aging.
Requirements of	 The candidate should have a MSc and experience with molecular and cellular biology. Our lab effort the RhD candidate state of the art equipment and expertise to address the scientific questions stated.
candidate:	 Our lab offers the PhD candidate state-of-the-art equipment and expertise to address the scientific questions stated above. Our lab consists of a mix of national and international PhD students and Postdocs and has an infrastructure
	that ensures intensive supervision and training during the PhD program.
	• Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the
	 scientific part of your scholarship proposal) English language requirement: IELTS 7.0(<i>min 6.0 for all subs</i>), TOEFL 100(<i>min 20 for all subs</i>)

School/Department:	Molecular Genetics Department, Erasmus MC
Supervisor information:	 Prof. Dr. Jurgen Marteijn (Full Professor on Transcription Stress and DNA damage response) J.Marteijn@erasmusmc.nl www.genomestability.nl
<u>Sciences</u>	 Grants and Prizes: 2019: AMMODO Science award for groundbreaking research (€1.200.000) 2019: VICI Grant of Netherlands Organization for Scientific Research (€1.500.000). 2014: VIDI Grant of Netherlands Organization for Scientific Research (€800.000). 2011: Erasmus MC Fellowship (€ 400.000).
	 5 Selected papers: 1: Elongation factor ELOF1 drives transcription-coupled repair and prevents genome instability. Geijer M,, Marteijn JA. Nature Cell Biology (Accepted 2021)
	2: The DNA damage response to transcription stress Lans H,, Marteijn JA Nature Reviews Molecular Cell Biology (2019)
	3: The core spliceosome as target and effector of non-canonical ATM signalling. Tresini M,, Marteijn JA. Nature (2015)
	4: Enhanced chromatin dynamics by FACT promotes transcriptional restart after UV-damage. Dinant C,, Marteijn JA Molecular Cell , (2013).
	5: UV-sensitive syndrome protein UVSSA recruits USP7 to regulate TCR. Schwertman P,, Marteijn JA. Nature Genetics (2012).
Project Title:	The molecular mechanism of DNA damage-induced aging
Abstract:	Due to the improved life span, age related diseases and discomfort have become a major social and medical issue. It is thus highly relevant to understand the biological processes that could counteract this phenomenon. Accumulation of DNA damage is a major contributor of age-related diseases. DNA damage blocks the transcription process, which is a crucial process for proper cell function. If the DNA damage that blocks transcription is not properly repaired it will result in cellular dysfunction, apoptosis and senescence, finally resulting in DNA damage induced aging. Cells counteract these deleterious effects by transcription-coupled repair (TCR), which removes the DNA damage thereby resolving the transcriptional block. The severe developmental problems and premature aging features of Cockayne syndrome patients - characterized by a hereditary TCR defect - underscore the importance of this process. Our lab is one of the world leading labs in the TCR field, and has recently identified several new repair factors in this pathway including UVSSA and ELOF1. Despite detailed knowledge on the TCR mechanism itself, surprisingly little is known about the last crucial step of TCR; how transcription restarts if the DNA damage is repaired. Using a multi-disciplinary approach of state-of-the-art live cell imaging and proteomic tools, the PhD student will study the molecular mechanism of transcription recovery after DNA repair. In addition, using unbiased CRISP/CAS9 based whole genome screens and advanced quantitative interaction proteomics studies we will identify novel proteins involved in this process. Together this will result in crucial new insights in TCR and will help to counteract the aging process.
Requirements of candidate:	 The candidate should have a Master and experience with molecular/cellular biology. Our lab offers the PhD candidate state-of-the-art equipment and expertise to address the scientific questions stated above. Our lab consists of a mix of both national and international PhD students and Post-docs and has an infrastructure that ensures intensive supervision during the PhD program. Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) English language requirement: IELTS 7.0(<i>min 6.0 for all subs</i>), TOEFL 100(<i>min 20 for all subs</i>)

School/Department:	Department of Molecular Genetics, Erasmus MC
Supervisor information:	Dr. Nitika Taneja, Ph.D., Principal Investigator and Group Leader
	• Email: n.taneja@erasmusmc.nl
World no 30 Biomedical	Website: <u>https://www.erasmusmc.nl/en/research/researchers/taneja-nitika</u>
Sciences	• Grants:
<u>Sciences</u>	Women in STEM Incentive grant by NWO, 2021
	• Erasmus+, 2020
	Young investigator award by Daniel den Hoed Stichting Fonds, 2018
	Most important publications:
	Lo et al. (2021) Science Advances PMID: 33952518
	• DiPiazza et al. (2021) PNAS PMID: 34035174
	Taneja et al. (2017) <i>Molecular Cell PMID:28318821</i>
	Taneja and Grewal (2017) <i>Cell Cycle PMID: 28805495</i>
	Mizuguchi et al. (2017) PNAS PMID: 28490498
	Mizuguchi et al. (2014) <i>Nature PMID: 25307058</i>
	Lee et al. (2013) <i>Cell PMID: 24210919</i> Development at al. (2012) <i>Plac Biology PMID: 22200276</i>
	Raychaudhuri et al. (2013) <i>Plos Biology PMID: 23300376</i>
Project Title:	Targeting chromatin modifiers for novel chemotherapeutic regimens
Abstract:	DNA replication is an essential but a precarious cellular process of central importance both to the development of
Abstruct.	cancer and its treatment. Indeed, failures in the replication process, for instance mutations in critical elements of
	the chromatin remodeling pathways, contribute to genome instability, an early event in tumorigenesis. The primary
	research goal of my lab is to obtain mechanistic understanding of pathways mediated by chromatin remodeling
	which allow stabilization of DNA replication machinery in normal as well as cancer cells. Such pathways play
	important role in in the hyper-proliferation of cancer cells and could also drive resistance towards chemotherapy.
	Therefore, chromatin modifying factors could become the potential candidates to be targeted for better therapies
	for the treatment of cancer as they are frequently mutated in cancerous cells but not in normal cells. We have
	recently identified a novel pathway and proteins involved in this pathway, which if targeted, can be exploited in the
	development of novel cancer therapeutic regimens.
	The focus of this project is to further understand the mechanistic link between chromatin remodeling pathways and
	the stability of DNA replication machinery to proper chromatin organization and concomitant genome stability.
	Through our research, we are trying to obtain a mechanistic understanding of the chromatin modifying (post- translational histone modifying) processes that render cells sensitive or resistant to commonly used
	chemotherapeutic treatments.
	Main methodology and techniques: The candidate will be part of a research team, including a senior postdoc as a
	daily supervisor, a PhD student working on a parallel project and a technician expert in sevaral techniques used in
	our lab. Our lab uses multidisciplinary approach combining high-thoughput genomics, quantitative imaging and high-
	thoughput proteomics. We use 2-D normal as well as human cancer cell lines and mouse 3-D tumor organoids for
	our studies. We frequently use CRISPR/Cas9 genome editing, Next generation sequencing analysis of chromatin via
	ChIP-Seq, 3-D chromatin organization via Hi-C, super-resolution imaging using SIM/STORM microscopes, single cell-
	based quantitative (QIBC) imaging and quantitative proteomics.
	PI:Nitika Taneja at ErasmusMC
	"Replication stress" Board of examiners, B.Sc/M.Sc Nanobiology program
	Our group (pre-Covid picture) Teacher at Erasmus MC & TU-Delft
Requirements of	We are looking for a highly motivated, hardworking student with master's degree to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills. English
candidate:	requirements: English speaking countries & Netherlands: no requirement
	Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)
	We offer: Supervision, lab facilities and infrastructure, and training. We will cover Laboratory costs. As a candidate PhD student at
	Erasmus MC, your salary and living expenses will be covered by your university or Scholarship Council. For more information
	regarding this vacancy, please contact n.taneja@erasmusmc.nl.

School/Department:	Molecular Genetics Department, Erasmus MC
Supervisor information:	Prof.Dr. W. Vermeulen and Dr. A. Pines
erc World no 30 Biomedical Sciences	 w.vermeulen@erasmusmc.nl and a.pines@erasmusmc.nl
	• <u>www.vermeulenlab.com</u>
Biomedical Sciences	• Grants and Prizes (selected):
	- Oncode Institute, Principle Investigator (2017); - Worldwide Cancer Research Project Grants (2015, & 2017); - Dutch Cancer Society (KWF), Research Grants (2016, & 2017); - European Research Council, ERC Advanced Grant (2013); - Dutch Scientific Organization, NWO-ENW-TOP grant (2018)
	• 5 Selected papers:
	 Ubiquitin and TFIIH-stimulated DDB2 dissociation drives DNA damage handover in nucleotide excision repair. Ribeiro-Silva C, Vermeulen W (corr. Auth.), and Lans H. Nature Commun.(2020). The DNA damage response to transcription stress. Lans, H., Hoeijmakers, J., Vermeulen, W*. and Marteijn, J.A*. (*corr. Auth.) Nature Rev.Mol.Cell.Biol. (2019) DNA damage sensitivity of SWI/SNF-deficient cells depends on TFIIH subunit p62/GTF2H1. Ribeiro-Silva, C.,, Vermeulen, W. Nature Commun. (2018). TRiC controls transcription resumption after UV damage by regulating Cockayne Syndrome protein A. Pines, A., Vermeulen, W.*, Pannu, N.S.* and Attikum, H.* (*corr. Auth.) Nature Commun. (2018). The core spliceosome as target and effector of non-canonical ATM signalling. Tresini M,, Vermeulen W.(corr.Auth.) Marteijn JA. Nature (2015).
Project Title:	Transcription stress: a link between DNA damage and aging
Abstract:	DNA is continuously damaged by environmental pollutants, radiation, and common cellular metabolites. DNA lesions interfere with genomic function, including transcription.
	Transcription-blocking lesions are removed by Transcription-Coupled Nucleotide Excision Repair (TC-NER), initiated by lesion-stalled RNApolymerase and subsequent binding of the
	 Cockayne Syndrome (CS) A and B proteins. Inherited <i>CSA</i> and <i>CSB</i> mutations are associated with serious health threats; including accelerated aging, developmental arrest and progressive neurodegeneration. Our research is aimed to provide mechanistic insight into the functional crosstalk between TC-NER-deficiency, DNA damage signaling, gene expression, and protein homeostasis by applying a multi-disciplinary approach combining innovative state-of-the-art technologies. To investigate the cell-specific consequences of <i>CSA</i> and <i>CSB</i> mutations, we will use CRISPR/CAS9-mediated gene editing combined with induced pluripotent stem cells (iPSC) reprogramming and cell-specific differentiation. The different cells will be used for quantitative mass-spectrometry to reveal the dynamic TC-NER interactome; RNA-sequencing to monitor transcription stress; live cell imaging to follow protein dynamics; super-resolution microscopy and biochemical 'protein aggregation' assays to study the protein homeostasis. The PhD student will participate in this frontline ambitious project aimed to obtain important mechanistic insight into the functional significance of TC-NER to counteract general DNA damage-induced diseases, including the molecular basis of neurodegeneration. Our lab offers: - state-of-the-art equipment and expertise to address the scientific questions stated above an internationally oriented work environment excellent PhD-training and coaching ensured through established Institutional and Departmental training and supervision programs.
Requirements of	 We are looking for highly motivated students that have a Master and thorough knowledge of molecular and cellular biology. English language requirement:
candidate:	English speaking countries & Netherlands: no requirement
	✓ Other countries: IELTS 7.0(min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Neuroscience Erasmus MC
Supervisor information:	Dr. Aleksandra Badura (Associate Professor)
	• Email: <u>a.badura@erasmusmc.nl</u> Website: <u>https://neuro.nl/research/badura</u>
World no 30 Biomedical	Grants:
Sciences World no 48 Neuroscience & Behavior	 Horizon 2020, Marie Sklodowska Curie Actions Innovative Training Network (PIPgen <u>https://pipgen.eu/</u>) Dutch Research Council (NWO) Starting Grant Vidi Dutch Research Council (NWO) Postdoctoral Fellowship Veni Erasmus MC Pilot grant
	Most important publications:
	 Badura A., Verpeut J.L., Metzger J.W, Pereira T.D, Pisano T.J., Deverett B., Bakshinskaya D.E., Wang S.SH. Normal cognitive and social development require posterior cerebellar activity. eLife 2018; 7, e36401. Giovannucci A.*, Badura A.*, Deverett B., Najafi F., Pereira T.D., Gao Z., Ozden I., Kloth A.D., Pnevmatikakis E., Paninski L., De Zeeuw C.I., Medina J.F., Wang S.SH. Cerebellar granule cells acquire a widespread predictive feedback signal during motor learning Nature Neurosci. 2017; 20, 727–734. Wang S.SH, Kloth A.D., Badura A. The Cerebellum, Sensitive Periods, and Autism. Neuron 2014; 83 (3), 518- 532. Badura A. *, Schonewille M. *, Voges K., Galliano E., Renier N., Gao Z., Witter L., Hoebeek F.E., Chédotal and De Zeeuw C.I. Climbing fiber input shapes reciprocity of Purkinje cell firing. Neuron 2013; 78, 700-13. Wulff P., Schonewille M., Renzi M., Viltono L., Sassoè-Pognetto M., Badura A., Gao Z., Hoebeek F.E., van Dorp S., Wisden W., Farrant M., De Zeeuw C.I. Synaptic inhibition of Purkinje cells mediates consolidation of vestibulo-
	cerebellar motor learning. Nature Neurosci. 12, 2009 1042-9.
Project Title:	Functional role of a novel ASD risk gene in the developing and adult brain
Abstract:	Genetic studies have implicated our gene of interest as a candidate gene for autism-spectrum disorder (ASD); however, a causal relationship between this gene and ASD does not exist. Recently, we identified a patient with biallelic mutations in this gene that presented with ASD, poor motor skills, intellectual disability, and hyperactivity. To fully understand the underlying pathology, we generated a mouse model with the patient-specific mutations. The mutant mice displayed gross impairments in motor coordination and sensorimotor learning as well as ASD-related behavioral abnormalities, hyperactivity, and cognitive deficits. We found that the patient and the mouse model show cerebellar anatomy and hypoplasia of several midbrain regions. We established that this gene is expressed in GABAergic neurons within the substantia nigra (SN) and ventral tegmental area (VTA) where mutant mice show a dramatic loss of GABAergic cells. The aim of this project is to answer the following questions: (1) How does the novel ASD risk gene regulate cerebellar development and how does its deficiency affect cerebellar functioning? (2) Which behavioral phenotypes are affected by the loss of GABAergic cells in the SN and VTA?
Requirements of	• We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using
candidate:	 teamwork to tackle important scientific questions and thus requires a student with good communication skills. Master degree in biochemistry, biophysics, neuroscience, or life sciences. Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) Proficiency in at least one of the coding languages: MATLAB, Python, C, Java, C++ Biomedical skills: Experience with Western blot, qPCR, PCR is required. Previous experience with mouse experiments is not a prerequisite but is welcomed. Neuroscience skills: General histology and immunocytochemistry. Candidates with experience in optogenetics or electrophysiology will be given a preference. English language requirement:
	 English speaking countries & Netherlands: no requirement

Dr. J. Gerard G. Borst, Professor of Neurophysiology (promotor)
il: g.borst@erasmusmc.nl
site: www.neuro.nl
onal Grants:
IMW-TOP 2018 (665 k€)
MSCA-ITN-2016 (total 2.5 M€)
ch Scientific Organization (ALW-Open) Grant, 2013, 2015 (300 k€ each)
ro-Basic Pharma Phenomics (FES0908) (2010; total 13 M€)
t important publications:
ure 383, 431-434 (1996)
uron 23, 821-832 (1999);
nce 289, 953-7 (2000);
nce 327: 1614-1618 (2010);
ure Neurosci. 13: 1050-1052 (2010);
Rev Physiol. 74:199-224 (2012);
ron 78: 936-948 (2013);
S 114: 4249-4254 (2017);
eurosci. 38: 2057-2068 (2018).
e 8, doi: 10.7554/eLife.49091 (2019).
nal mechanisms underlying tinnitus
s is a very common disorder in which a patient hears sound in the absence of an
I source. Severe tinnitus can have a devastating impact on the quality of life, but
the large burden of disease there is currently no curative treatment, and the mainstay
py currently focusses on helping patients cope with their tinnitus. A substantial
ck in developing an effective treatment for tinnitus is the lack of understanding of the
athological mechanisms underlying it.
project you will investigate the cellular mechanisms underlying tinnitus. To test this, you
estigate in mice whether cortical feedback inhibition is altered in the inferior colliculus
als with tinnitus. The presence of tinnitus will be assessed by a novel operant
oning task, while neuronal IC activity and cortical feedback will be measured and
lated using in vivo optical (two-photon imaging, optogenetics) and electrophysiological
electrode; patch clamp) techniques. These experiments will provide novel insight into
mechanisms at both a cellular level and at the level of individual auditory regions,
vill constitute an important synergistic step towards the development of a curative
ent.
ire looking for a highly motivated student with interests in hearing research and preferentially experience with in vivo rdings to join our international team.
er degree or MD with research experience.
larship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the
tific part of your scholarship proposal). sh language requirement:
sn language requirement: English speaking countries & Netherlands: no requirement
Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Neuroscience Erasmus MC
Supervisor information:	Dr. P.A. Forbes, PhD and Prof. M.A. Frens
	• Email: p.forbes@erasmusmc.nl; m.frens@erasmusmc.nl; http://www.neuro.nl
World no 30 Biomedical	Personal Grants:
<u>Sciences</u>	 Dutch Scientific Organization Grant (VIDI, Top Talent, VENI), 2017, 2019, 2021 ESA Parabolic Flight Campaigns, 2016, 2017, 2018
Sciences	 EsA Parabolic Flight Campuigns, 2010, 2017, 2018 European Research Commission (Marie Sklodowska-Curie Action), 2014
World no 48 Neuroscience &	- National Science and Engineering Research Council (Canada), 2013
Behavior	- Nissan Motors, 2013
	Most important publications: alife 2021 doi: 10.7EE4/alife 6E08E
	 eLife, 2021, doi: 10.7554/eLife.65085 Scientific Reports, 2021, doi: 10.1038/s41598-021-93037-7
	- Journal of Neuroscience, 2020, doi: 10.1523/JNEUROSCI.1463-19.2020
	- Annals of Neurology, 2020, doi: 10.1002/ana.25679
	- Nature Communications 2019, doi: 10.1038/s41467-019-09738-1
	- Journal of Physiology, 2019, doi: 10.1113/JP278642
	 Frontiers in Physiology, 2019, doi: 10.3389/fphys.2019.00476 eNeuro, 2018, doi: 10.1523/ENEURO.0170-18.2018
	 Handbook of Clinical Neurology, 2018, doi: 10.1016/B978-0-444-63916-5.00004-5
	- Journal of Physiology, 2017, doi: 10.1113/JP272614
	 Journal of Neuroscience, 2016, doi: 0.1523/JNEUROSCI.1902-16.2016
Project Title:	Neuromechanical principles underlying the multiaxial control of human balance
Abstract:	Upright balance is a continuous struggle against Earth's gravitational pull. Our vertical posture
	is inherently unstable and must be balanced within a small base of support. Any difficulties in
	maintaining upright balance puts us at risk of serious injuries due to falls, bringing personal,
	societal and economic burdens that will continue to increase without a comprehensive
	understanding of the mechanisms underpinning standing balance. Ongoing balance control
	relies on complex interactions between our body's biomechanics and the neural (sensory, motor
	and cognition) systems contributing to standing. For example, the brain must account for the
	fact the muscles generating torque around our joints often cross axes, meaning that any
	passive/active muscle tension influences joint torques in multiple directions (i.e. cross-talk).
	While these biomechanical and neural factors of balance have intrigued researchers for
	decades, methodological difficulties in unraveling their interactions provides an incomplete
	picture of how the brain controls standing. The long-term aim of our research is to disentangle
	these biomechanical and neural contributions to standing balance by combining robotic
	simulation, human neurophysiology (EEG/EMG), computational modeling and sensory
	stimulation to push the field passed these obstacles. This project will determine how
	biomechanical and neural factors along our two primary axes of balance are coordinated to
	maintain balance, establishing whether cross-talk between their control impedes or enhances
	our adaptation to the daily challenges of balance. In addition, this project will reveal how
	sensory and motor cues of balancing self-motion govern the conscious perception and control
	during imposed sensorimotor errors. Finally, by performing experiments in healthy participants
	and patients (i.e. vestibular loss and cerebellar ataxia), we will directly test how disruption at
	different levels of balance influence the brain's ability to adapt and learn. Overall, this innovative
	research will reveal causal relationships between the neural computations and compensatory
	responses required for balance and locomotion.
Requirements of	• We are looking for a highly motivated student with interests in hearing research and preferentially experience with in vivo
candidate:	recordings to join our international team.
cananaace.	 Master degree or MD with research experience. Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the
	scientific part of your scholarship proposal).
	English language requirement:
	English speaking countries & Netherlands: no requirement
	• Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Neuroscience Erasmus MC
Supervisor information:	• dr Johan JM Pel, associate professor
	• Email: j.pel@erasmusmc.nl
World no 30 Biomedical Sciences	Website: http://www.neuro.nl/research.php
World no 48 Neuroscience &	• Personal Grants:
Behavior	- ZonMW grant 2009, 2012, 2018
<u>Benavior</u>	- Zon MW – DST India grant 2012
	 Most important publications:
	- Transl Vis Sci Technol. 2019 Jul 30;8(4):13.
	- Graefes Arch Clin Exp Ophthalmol. 2019 Apr 3
	- Brain Dev. 2018 Oct 6. pii: S0387-7604(18)30469-8.
	 Cerebellum. 2018 Sep 14. doi: 10.1007/s12311-018-0975-9 Graefes Arch Clin Exp Ophthalmol. 2018 Feb;256(2):371-379
	- J Vis. 2016;16(5):18
	- Dev Med Child Neurol. 2016 Oct;58(10):1030-5
	- Motor Control. 2016 Jan;20(1):1-20
	- J Vis Exp. 2016 Jul 9;(113)
	- J Ophthalmol. 2015;2015:425067
	- J Parkinsons Dis. 2014 4:599–608
	- Invest Ophthalmol Vis Sci. 2013 Mar 5;54(3):1656-64
	- J Alzheimers Dis. 2012 Jan 1;30(1):131-43
Project Title:	Visual-motor and visual vestibular interactions
Abstract:	The reflex movements that we display as a baby gradually develop into complex goal-
	directed behavior, which is essential for development and learning. The underlying
	sensorimotor integration translates visual, vestibular and somatosensory information
	into (in)voluntary motor output during complex behaviors such as standing balance or
	goal-directed arm movements. In children, abnormal performance scores of
	neuropsychological and motor tests signal integration problems. They fail, however, in
	revealing which underlying functions, e.g. visual, motor or visuomotor integration, are
	impaired. In elderly, neurodegeneration may result in deficits in the sensorimotor
	integration network leading to behavioral problems. In our group, we are interested in
	the fundamental and clinical relevance of quantitatively assessed (altered) eye, hand
	and body movements during sensorimotor integration tests. To achieve this goal, we
	develop new techniques, including advanced eye movement recordings (imprinted
	lenses) and combine them with quantitative assessment of visuomotor integration
	performances and interactions. Ultimately, our approaches allow us to determine how
	different sensory modalities interact and how they contribute to the development and
Doguiromonte of any didate	 control of motor and non-motor functions. We are looking for a highly motivated, hardworking student to join our international team. Our strength is to
Requirements of candidate:	tackle large scientific questions and thus requires a student with good communication skills.
	Master degree or MD
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal)
	 English language requirement:
	• English speaking countries & Netherlands: no requirement
	 Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Neuroscience Erasmus MC
Supervisor	Dr. Martijn Schonewille, <u>m.schonewille@erasmusmc.nl</u>
information:	https://neuro.nl/research/schonewille
	Personal Grants:
World no 30 Biomedical	- ERC Starting Grant (ERC-Stg), 2015
<u>Sciences</u>	- Dutch Scientific Organization (ALW-Open) Grant, 2014 (co-appl.)
	 Dutch Scientific Organization (ALW-Veni) Grant, 2011 Erasmus University Fellowship, EUR, 2010
World no 48 Neuroscience	- Grants for group members:
<u>& Behavior</u>	- Dutch Scientific Organization (ALW-Veni) Grant, 2018
	- German Research Organization (DFG) Grant, 2019
	- Dutch Scientific Organization (Offroad), 2020
	 South African Research Organization (NRF-Nuffic), 2020 Erasmus MC Fellowship 2021
	- Dutch Scientific Organization, NWO-XS, 2021 (2x)
	Most important publications:
	Nat Neurosci. 9(4):459-61; Neuron. 12;58(5):655-8; Nat Neurosci. 12(8):1042-9; Neuron. 26;67(4):618-28;
	Neuron. 14;70(1):43-50.; Nat Rev Neurosci. 12(6):327-44. Review; EMBO J. 7;31(5):1217-30; Neuron
	22;78(4):700-13; eLife; 10.7554/eLife.02536; Nat Commun. 2016 Sep 1;7:12627; PNAS 2021 September 7,
	2021 118 (36) e2016969118; eLife; 10.7554/eLife.45590.001; PNAS 2021 September 14, 118 (37) e2102635118; Nat Comm. 2021 12, Art#: 4129 (2021); eLife 2021;10:e63668;
Project Title:	Cerebellar differentiation in development of motor functions and
	neurodevelopmental disorders
Abstract:	The perfect execution of a voluntary movement requires the appropriate integration of current
	bodily state, sensory input and desired outcome. To assure that this motor output becomes and
	remains appropriate, the brain needs to learn from the result of previous outputs. The
	cerebellum plays a central role in sensorimotor integration, yet -despite decades of studies-
	there is no generally excepted theory for cerebellar functioning. We recently demonstrated
	that cerebellar modules, identified based on anatomical connectivity and gene expression,
	differ distinctly in spike activity properties. It is the lab's long-term goal to identify the ontogeny
	of anatomical and physiological differences between modules, and their functional
	consequences.
	To achieve this goal, we make use a variety of techniques including molecular approaches, in
	vitro and in vivo electrophysiology, 1p and 2p imaging techniques, optogenetic stimulation and
	behavioral evaluations. We aim to determine how differential gene expression patterns control
	the development of distinct physiological properties and anatomical connection patterns of the
	types of neurons in different cerebellar modules. We will determine the impact of the genetic
	differentiation in cerebellar input, processing and output.
	Ultimately, the combined results of these studies will reveal how distinct differences between
	cerebellar modules develop, and how the modular ensemble ensures proper cerebellar
	information processing for optimal coordination of timing and force of movements. Combined
	with the growing body of evidence for a cerebellar role in higher order brain functions and
	neurodevelopmental disorders, this knowledge will be fundamental for understanding how the
	juvenile brain develops.
Requirements of	• We are looking for a highly motivated, hardworking student to join our international team. Since we are tackling
candidate:	complex scientific questions regarding decision making, procedural learning, as well as memory disorders, we hope to find a student is willing to learn new techniques, has affinity with quantitative data analysis, and can communicate well.
	Master degree in (bio)physics or neuroscience, an engineering degree, or an MD.
	Scholarship that will cover subsistence allowance and international air plane ticket.
	• English language requirement: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs). When writing the CSC
	proposal we will help with the scientific part of your scholarship proposal.

School/Department:	Department of Neuroscience Erasmus MC
Supervisor	• Dr. Zhenyu Gao, z.gao@erasmusmc.nl; https://neuro.nl/research/gao
information:	Personal Grants:
•	- ERC Starting Grant (ERC-Stg), 2019
World no 30 Biomedical	- Dutch Scientific Organization (NWO-VIDI) Grant, 2019
<u>Sciences</u>	- Dutch Scientific Organization (NWO-Klein) Grant, 2019
Sciences	- Dutch Scientific Organization (NWO-CAS) Grant, 2017
World no 48 Neuroscience	- Erasmus MC Fellowship, 2016
<u>& Behavior</u>	- Dutch Scientific Organization (NWO-VENI) Grant, 2014
	 Most important publications: Nature 2018 563(7729):113-116
	- Elife 2017 15;6 pii:e28132
	- Neuron 2016 89(3):645-57
	- Cell Reports 2013 253(4):1239-51
	- Nature Reviews Neuroscience 2012 13: 619–635
	- Journal of Neuroscience 2012 31;32(44):15533-46
	- Neuron 2011 14;70(1):43-50
Project Title:	Dissecting the brain-wide connectome for motor planning
Abstract:	All voluntary movements are directed by proper motor plans in the brain. How does the brain
	effectively generate these motor plans and use them to direct future movements? Previous
	studies suggested that the motor cortex play a key role in motor planning. Motor cortical
	neurons maintain their activity for seconds before the movement's onset, which allows the
	brain to temporarily retain valuable information to secure accurate execution of the motor
	plans. Our recent research provided evidence for the functional involvement of the cerebellum
	in motor planning (Gao <i>et al</i> , Nature 2018). For this PhD project we will focus on further
	dissecting the brain-wide circuits that are relevant for motor planning. We will examine
	whether the sensorimotor representation from the cerebral cortex is integrated in cerebellum
	during motor planning and that the computation in cerebro-cerebellar circuits is instrumental
	for supporting the preparatory activity. We will use an integrative approach to 1). identify the
	cerebrum-to-cerebellum inputs that are relevant for motor planning; 2). determine how
	cerebellar circuits integrate cerebral inputs and generate corresponding outputs during motor
	planning; 3). Identify the role of cerebellar outputs in motor planning and explore their
	computational mechanisms. This project will greatly advance our knowledge on the general
	computational principles underlying motor planning. In the future it will pave the way to a
	mechanistic understanding of brain-wide communication in cognitive tasks with its influence
	extended to future computer science, humanized prosthetics, and medicine.
Requirements of	We look for highly motivated students to join our multi-disciplinary team. We welcome students with Msc in
	biotechnology, neuroscience, bio-engineering, and other life sciences majors. Prior experience in molecular biology,
candidate:	imaging, electrophysiology and computational modelling is preferred, but not essential.
	 Master degree in (bio)physics or neuroscience, an engineering degree, or an MD. Scholarship that will cover subsistence allowance and international air plane ticket
	Scholarship that will cover subsistence and wance and international an plane taket.
	 English language requirement: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs). When writing the CSC
	proposal we will help with the scientific part of your scholarship proposal.

School/Department:	Department of Neuroscience Erasmus MC
Supervisor	Prof. Dr. Chris I. De Zeeuw, <u>c.dezeeuw@erasmusmc.nl</u>
information:	 https://neuro.nl/research/de-zeeuw
	Personal Grants:
World no 30 Biomedical	- ERC Advanced Grant (ERC-Adv), 2014
<u>Sciences</u>	- ERC PoC grants (ERC-PoC), 2015, 2016, 2017
	- Dutch Scientific Organization (ALW-Open) Grants, 2016, 2017
World no 48 Neuroscience	- ZonMw Grant, 2016
<u>& Behavior</u>	- KNAW Grants, 2017, 2018
	 Most important publications:
	- <u>Nature Neuroscience 2021 24: 160</u> - <u>Nature Reviews Neuroscience 2021 22:92</u>
	- Nature Communications 2020 11 - Nature Communications 2019 10
	- Nature 2018 563:113 - Nature Communications 2018 9
	- Science Adv 2018 4 - Science 2017 356:1084 - Nature Neuroscience 2017 20:727 - Neuron 2017 93:409
Project Title:	Cerebro-cerebellar Interactions during Cognitive Processing
Abstract:	Coordinating cognitive processes forms the most important and complex task of the brain. Not surprisingly, coordinated control of these functions requires intensive communication within and between many brain regions. Of crucial importance is the mutual communication between cerebellum and cerebral cortex (De Zeeuw, 2021, <i>Nature Reviews Neuroscience</i> ; Gao et al., 2018, <i>Nature</i>). This becomes apparent, for instance, in patients suffering from autism (Peter et al., 2016, <i>Nature Commun</i>), spino-cerebellar ataxia (Hoogland et al., 2015, <i>Current Biol</i>), or Alzheimer's (Sepulveda-Falla et al., 2014, <i>J. Clin. Invest</i> .), in which the output neurons of cerebellum and cerebral cortex become dysfunctional. Before we can start to understand such pathology, we need to comprehend cerebello-cerebral communication under the normal conditions, like decision making and motor planning. For this reason we have developed a behavioral paradigm in which mice are being trained to use their whiskers to discriminate the location or properties of an object, to make a decision based on their sensory input during a delay period, and to report their decision as licking into a trained direction (Gao et al., 2018, <i>Nature</i>). This task has been shown to require proper functioning of the cerebellum and cerebral cortex, but it is unclear how subcortical structures ultimately determine direction encoding in this process (Boele et al., 2018, <i>Science Adv</i>). For this CSC project we will 1) record neuronal activity in the cerebellum, cerebral cortex and subcortical structures simultaneously in normal mice during and after training; 2) selectively modulate neuronal activity during and after training using optogenetics; and 3) rescue phenotypes in mouse models of autism, ataxia and Alzheimer's. Together, these specific aims should allow us to elucidate how interactions between cerebellum and cerebral cortex drive complex cognitive and motor tasks, and
Requirements of candidate:	 compensate for dysfunctions thereof in wide-spread brain diseases. We are looking for a highly motivated, hardworking student to join our international team. Since we are tackling complex scientific questions regarding decision making, procedural learning, as well as memory disorders, we hope to find a student is willing to learn new techniques, has affinity with quantitative data analysis, and can communicate well. Master degree in (bio)physics or neuroscience, an engineering degree, or an MD. Scholarship that will cover subsistence allowance and international air plane ticket. English language requirement: IELTS 7.0 (<i>min 6.0 for all subs</i>), TOEFL 100 (<i>min 20 for all subs</i>). When writing the CSC
	proposal we will help with the scientific part of your scholarship proposal.

Dept. of Oral & Maxillofacial Surgery, Special Dental Care & Orthodontics

School/Department:	Department of Oral & Maxillofacial Surgery, Special Dental Care & Orthodontics
	Erasmus MC
Supervisor information:	Prof dr Eppo Wolvius – Head of Department Prof dr. Fernando Rivadeneira Email: <u>e.wolvius@erasmusmc.nl</u> <u>f.rivadeneira@erasmusmc.nl</u> Website: <u>https://www.oral-health.nl/</u> Grants: Europagen Reference Naturack on Crapic disagrees https://arn.grapic.org
world no 13 Surgery world no 36 in Radiology, Nuclear Medicine and Imaging	 European Reference Network on Cranial diseases https://ern-cranio.eu European Commission Cost Action: GEnomics of MusculoSkeletal traits TranslatiOnal Network (CA86139) https://www.cost.eu/actions/CA18139/ European Commission MSC-ITN Tissue engineering in osteoarthritis and bone disease https://www.carbonresearch.eu. ERC Advanced grant 2021 Most important publications: Vucic, S., R. W. Drost, A. J. van Wijk, P. R. Wesselink and E. B. Wolvius (2016). "Patterns of orodental injury and mouthguard use in Dutch field hockey." Br J Sports Med 50(11): 661-668. Vucic, S., R. W. Drost, E. M. Ongkosuwito and E. B. Wolvius (2016). "Dentofacial trauma and players' attitude towards mouthguard use in field hockey: a systematic review and meta-analysis." Br J Sports Med 50(5): 298-304. Jonsson, L., T. E. Magnusson, A. Thordarson, T. Jonsson, F. Geller, B. Feenstra, M. Melbye, E. A. Nohr, S. Vucic, B. Dhamo, F. Rivadeneira, E. M. Ongkosuwito, E. B. Wolvius, E. J. Leslie, M. L. Marazita, B. J. Howe, L. M. Moreno Uribe, I. Alonso, M. Santos, T. Pinho, R. Jonsson, G. Audolfsson, L. Gudmundsson, M. S. Nawaz, S. Olafsson, O. Gustafsson, A. Ingason, U. Unnsteinsdottir, G. Bjornsdottir, G. B. Walters, M. Zervas, A. Oddsson, D. F. Gubbjartsson, S. Steinberg, H. Stefansson and K. Stefansson (2018). "Rare and Common Variants Conferring Risk of Tooth Agenesis." J Dent Res 97(5): 515-522. Vucic, S., T. I. M. Korevaar, B. Dhamo, V. W. V. Jaddoe, R. P. Peeters, E. B. Wolvius and E. M. Ongkosuwito (2017). "Thyroid Function during Early Life and Dental Development." J Dent Res 96(9): 1020-1026. Alslanaj, B., L. Kragt, I. Voshol, M. Koudstaal, M. A. Kuijpers, T. Xi, S. J. Berge, C. Vermeij-Keers and E. M. Ongkosuwito (2017). "Dentition Patterns in Different Unilateral Cleft Lip Subphenotypes." J Dent Res 96(13): 1482-1489 Liu, X., Kayser, M., Ku
Project Title:	Three-dimensional (3D) Facial Shape Analysis using Artificial Intelligence
Abstract:	The human face is complex three-dimensional structure that makes each of us uniquely distinguishable, but strongly determined by genetic factors. Consequently, many developmental, psychiatric and genetic abnormalities have defined facial morphological features. However, the underlying complexity of facial morphology cannot be fully captured by simple geometric measures. Rather, it is now increasingly clear that the genetic determination of facial morphology and its relation with health outcomes requires more sophisticated quantitative approaches for capturing facial morphology. Recent advances in computational and methodological approaches have made possible accurate and precise derivation of facial traits. This project will focus on developing methods (based on machine learning and deep learning technologies) to derive complex facial measurements. the ultimate aim of this project is to leverage the large-scale 3D facial imaging, which provides extensive genetic and epidemiological measures, to unravel the complexity between genetics, facial morphology
	and health outcomes.
Requirements of candidate:	 We are looking for a highly motivated, hardworking student to join our very international team. Successful candidates are expected to have a strong quantitative or computer science background, excel at critical thinking, with strong motivation to engage in development and application of advanced analytical methods. Master degree in mathematics, computer science, statistics, bioinformatics, physics, electrical engineering, or in an equivalent discipline. Experience with: Python, linux, shell. Experience with machine learning methods. deep learning methods is advantage Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we can help with the scientific part of your scholarship proposal)
	 English language requirement: English speaking countries & Netherlands: no requirement; Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

Dept. of Oral & Maxillofacial Surgery, Special Dental Care & Orthodontics

School/Department:	Department of Oral & Maxillofacial Surgery, Special Dental Care & Orthodontics
	Erasmus MC
Supervisor information:	Prof. Eppo Wolvius (<u>e.wolvius@erasmusmc.nl</u>), Head of the Department
world no 13 Surgery	Prof. Fernando Rivadeneira (f.rivadeneira@erasmusmc.nl), Full Professor
	Dr. Lea Kragt (<u>I.kragt@erasmusmc.nl</u>), Post-doctoral Scholar
	Website: www.oral-health.nl
	Most important publications:
	2016: J Dent Res 95(4):395-401.
	2016: Caries Res 50(5):471-479 & 489-497
	2017: J Dent Res 96(13): 1482-1489.
	2017: J Dent 62:18-24.
	2018: Hum Mol Genet 27(17):3113-3127.
	2019: Qual Life Res 28(7):1783-1791.
	2020: Bone 132:115-180. 2021: J Nutr. 151(7):1993-2000
Project Title:	The oral microbiome in adolescents - individual, environmental and genetic
Floject Inte.	determinants
Abstract	The department of oral and maxillofacial surgery, special dental care and orthodontics
Abstract:	conducts oral health research in big datasets from population-based cohorts and clinical
	cohorts. Oral health research in this setting is worldwide nearly unique. Dr Lea Kragt has
	worked within this research line for 8 years, is coordinating the collection of dental data and
	has initiated and conducted research on different aspects within the research group, from
	quality of life factors to endocrine disrupters. We offer an interesting and challenging position
	in an ambitious yet friendly scientific and clinical research environment.
	PhD project:
	The oral microbiome offers an innovative approach to develop new preventive strategies for
	dental diseases. Dental caries for example is a major public health problem with a prevalence
	around 30% in Dutch children and up to 90% among children worldwide, typically affecting in
	larger proportions socially disadvantaged and marginalized populations. Though caries is a
	preventable disease, due to its multifactorial nature, the condition is difficult to tackle.
	Therefore the aim of this project is to provide a basis for the use of the oral microbiome in
	both risk-identification and progression-control of dental caries by understanding its
	composition and modifiability. Dental biofilm samples have been collected (n=4800) and are
	processed using 16S rRNA sequencing to obtain oral microbiome profiles. Logistic regression
	(alpha diversity) and permutation analysis (beta diversity) will be used to identify associations
	between general as well as oral health factors and oral microbiome profiles. The candidate for
	this project is free to develop additional research objectives related to the oral microbiome
	during the project.
Requirements of	• We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in
candidate:	 using team work to tackle large scientific questions and thus requires a student with good communication skills. Research Master degree (epidemiology, biomedical, (micro)biology or equivalent) or doctor of medicine (MD) or
	doctor of dentistry (DD) required
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal)
	 scientific part of your scholarship proposal) English language requirement:
	English speaking countries & Netherlands: no requirement
	Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

Dept. of Oral & Maxillofacial Surgery, Special Dental Care & Orthodontics

School/Department:	Department of oral and maxillofacial surgery, special dental care and orthodontics,
	Erasmus MC
Supervisor information:	Prof. Eppo Wolvius (<u>e.wolvius@erasmusmc.nl</u>), Head of the Department
	Prof. Fernando Rivadeneira (<u>f.rivadeneira@erasmusmc.nl</u>), Full Professor
erc	Dr. Lea Kragt (<u>l.kragt@erasmusmc.nl</u>), Post-doctoral Scholar
world no 13 Surgery	Website: <u>www.oral-health.nl</u>
	Most important publications:
	2016: J Dent Res 95(4):395-401.
	2016: Caries Res 50(5):471-479 & 489-497 2017: J Dent Res 96(13): 1482-1489.
	2017: J Dent 62:18-24.
	2018: Hum Mol Genet 27(17):3113-3127.
	2019: Qual Life Res 28(7):1783-1791.
	2020: Bone 132:115-180.
Project Title:	Oral health trajectories - individual, environmental and genetic determinants
Abstract:	The department of oral and maxillofacial surgery, special dental care and orthodontics
	conducts oral health research in big datasets from population-based cohorts and clinical
	cohorts. Oral health research in this setting is worldwide nearly unique. Dr Lea Kragt has
	worked within this research line for 8 years, is coordinating the collection of dental data and
	has initiated and conducted research on different aspects within the research group, from
	quality of life factors to endocrine disrupters. We offer an interesting and challenging position
	in an ambitious yet friendly scientific and clinical research environment.
	PhD project:
	Dental caries is a major public health problem with a prevalence around 30% in Dutch
	children and up to 90% among children worldwide. Next to this, dental caries is socially
	patterned, typically affecting in larger proportions socially disadvantaged and marginalized
	populations. The disparities already exist early in childhood, but increase throughout the
	lifetime. Carious lesions are very common in children, but the transition from childhood to
	adulthood is an even more sensitive period for the development of oral health and disease.
	The underlying mechanisms in the association of disadvantaged populations with oral
	diseases are not clear.
	The candidate will identify and investigate distinct trajectories of oral health and disease in
	growing children/young adults using latent class models. Multinomial multilevel regression
	analysis will be performed to study the behavioral, environmental and genetic predictors of
	oral health trajectories. In addition, he/she will employ state of the art biomarkers (including
	genomic) assessments that provide additional insight to assess causal relationships between
	potentially confounded risk factors for oral diseases. For example, the potential role of the
	oral microbiome in the relation of individual and environmental factors and oral diseases
	might be explored considering a plausible mediation by these factors.
Requirements of	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills.
candidate:	 Research Master degree (public health, epidemiology or equivalent) or doctor of medicine (MD) or doctor of dentistry
	(DD)
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship property).
	 scientific part of your scholarship proposal) English language requirement:
	English speaking countries & Netherlands: no requirement
	Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

The Department of Pathology of the Erasmus Medical Center in Rotterdam, The Netherlands. <u>https://www.erasmusmc.nl/pathologie/research/?lang=en</u> Head of the Dept: Prof. Dr. F. van Kemenade.

In the Department of Pathology of the Erasmus MC the research topics can be grouped into two major themes: 1. Oncology and 2. Cardiovascular / transplantation-immunology. The cancer research is both translational and basal, and encompasses topics in cancers of the brain, urogenital and GI tract. In addition there are basic research topics in stem cell research and there is a Center for Optical Imaging in which various projects are being carried out.

Why choosing for this department?

The department of Pathology is well equipped with virtual all molecular techniques and a laboratory for molecular diagnostics is incorporated. The department harbors a accredited tissue bank of over 40,000 frozen specimens. In addition, being the largest department of pathology in the country there is a large FFPE archive, and a large archive of autopsy-related specimens. The department belongs to a cluster of service laboratories (Lab Medicine, Immunology, Microbiology, Radiology), but research collaborations are extending well beyond to departments of (clinical) genetics, experimental cardiology, nephrology / transplantation and more.

Key publications (2016-2017 of the senior PIs:)

Prof. Fodde (GI, stem cell biology): Schewe M et al., Cell Stem Cell. 2016.; Rodriguez-Colman MJ et al., Nature. 2017. Prof. Houtsmuller (Center for Optical Imaging): Sanchez H. Nucleic Acids Res. 2017; Meddens MB et al. Nat Commun. 2016.

Prof. Kros (Neuro-Onc) van den Bent MJ. et al. Lancet 2017; Zheng PP et al. Med Res Rev; 2017; Zhu C. et al. Neuro Oncol. 2017; Thompson EM et al. Lancet Oncol. 2016.

Dr. van Leenders (Urogenital) Roobol MJ et al. Eur Urol. 2017; Ruela-de-Sousa RR. et al. Eur Urol. 2016.; Alberts AR et al. Eur Urol. 2016.

Selected recent Honors & Awards:

Collaborative Grants (NWO, Horizon2020, MSCA, other):

NWO – Building blocks € 150K; KWF- Ovarian Cancer € 570K; KWF – Raman spectroscopy €635K; MLDS – Colon cancer € 240K; Horizon 2020 – SPIDIA4P € 119K; Industry – Roche €131K; Industry – Astrazenica €269K; Industry – MDX Health €578K.

School/Department:	Department of Pathology Erasmus MC
Supervisor information:	Prof dr Adriaan B. Houtsmuller Assoc. Prof dr Timo L.M. ten Hagen
	Email: <u>a.houtsmuller@erasmusmc.nl</u> <u>t.l.m.tenhagen@erasmusmc.nl</u>
world no 32 Oncology	Website: <u>www.erasmusmc.nl</u> , <u>www.molmed.nl</u> Create: <u>NWU_FU_FPG_FU_FPG_CSC_Mreas_NWC_</u> _PPOL_DdUSt
	 Grants: NIH, EU FP6, EU FP7, CSC, Mrace, NWO, BBOL, DdHSt Most important publications:
	1)ten Hagen TLM, Smits R, Bruno MJ, Fuhler GM, Peppelenbosch MP. Carcinogenesis. 2019 Feb 20
	2)ten Hagen TLM. Sci Rep. 2018 Jun 25;8(1):9596.
	 3)ten Hagen TLM,, Peppelenbosch MP, Fuhler GM. Oncotarget. 2016 8;7(45):73525-40. 4)ten Hagen TLM, Fuhler GM. Oncotarget. 2016 Apr 19;7(16):21922-38.
	5)ten Hagen TLM Nat Protoc. 2015 Jun;10(6):904-15.
	6)ten Hagen TL. Eur J Cancer. 2016 Jan;53:135-43.
	 7)Houtsmuller AB. Sci Rep. 2019 Jul 18;9(1):10460. 8)Houtsmuller AB, van den Dries K, Wiseman PW, Cambi A. Nat Commun. 2016 7:13127.
	9)Houtsmuller A, Huveneers S, de Rooij J. Sci Rep. 2015 5:17225.
	10)Houtsmuller AB, van de Water B. J Cell Sci. 2012 125(Pt 19):4498-506.
Project Title:	Understanding local and systemic progression of cancer with respect to tumor –
	stroma interaction and metastasis development.
Abstract:	Local development of cancer is not only interesting for development of therapeutics or
	understand what drives tumor progression. Importantly, aspects of local development connect
	with the occurrence of metastasis, progression of the disease and eventually mortality. For
	instance, while tumor cell proliferate and a larger mass is formed the surrounding tissue, tumor
	stroma, needs to be recruited. The environment (may) provide stimulatory signals, inflammatory
	cells promote growth, specific immune cells inhibit antitumor responses, nutrients and oxygen are
	delivered through a (newly) developed vascular bed. These all will help the tumor to progress
	locally. However, these factors as well affect progression beyond the primary tumor. Vasculature
	and lymphatics help metastasis by providing the logistics for spreading cells, inflammation may
	help cells to escape through opening tissues and endothelial lining, and locally produced factors
	may have an effect at distance, either by inhibiting or promoting growth of new tumors, or by
	creating a favorable niche at distance for circulating tumor cells to locate. It is clear that expansion
	of a tumor is not just a stochastic effect but that certain tumor cells are responsible for the onset of growth, which some would call tumor stem cells, and that expansion may involve a different set
	of tumor cells resulting from the stem cells. More so, when tumors evolve locally clonal growth
	may occur, but clearly differentiation of tumor cells takes place. For instance, it is proposed that
	cells go through transitions such as the EMT (epithelial-to-mesenchymal transition), where
	proliferation is tuned down and migratory capacity goes up when a cell is destined to metastasis.
	When at location this process is reversed; the tumor cells loses the migratory capacity while
	gaining again in proliferative capacity. However, we have examples where this is not a given;
	tumor cells exhibit high proliferation as well as migration capacities at the same time. Here we
	study the aspects of tumor progression as disease in a number of in vitro and in vivo models
	including, but not limited to, intravital microscopy, advanced 3D live cell imaging, spehriod
	cultures, clonal expansion, and vascular formation. Below 3D growth and dispersion in vitro (left
	two images) and intravital window with image of green vessels and red blood marker (right two
	images)
Requirements of	• We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in
candidate:	using team work to tackle large scientific questions and thus requires a student with good communication skills.
	 Master degree or MD Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the
	scientific part of your scholarship proposal)
	English language requirement: English engetting equatrics & Notherlands, no requirement
	English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Pathology Erasmus MC
Supervisor	Prof dr Adriaan B. Houtsmuller Assoc. Prof dr Timo L.M. ten Hagen
information:	Dr. Mohamadreza Amin
	Email: <u>a.houtsmuller@erasmusmc.nl</u> <u>t.l.m.tenhagen@erasmusmc.nl</u>
world no 32 Oncology	<u>M.amin@erasmusmc.nl</u> • Website: www.erasmusmc.nl , www.molmed.nl
	• Grants: NIH, EU FP6, EU FP7, CSC, Mrace, NWO, BBOL, DdHSt
	Most important publications:
	1-Seynhaeve, A.L.B et al. Hyperthermia and smart drug delivery systems for solid tumor therapy. Adv Drug Deliv Rev 2020. 2-Amin, M.; et al. Regulation of in vivo behavior of tat-modified liposome by associated protein corona and avidity to tumor cells. Int J Nanomedicine 2018,
	13, 7441-7455.
	3-Seynhaeve, A.L. et al Intact doxil is taken up intracellularly and released doxorubicin sequesters in the lysosome: Evaluated by in vitro/in vivo live cell imaging. J Control Release 2013, 172, 330-340.
	4-Li, L. et al. Improved intratumoral nanoparticle extravasation and penetration by mild hyperthermia. J Control Release 2013, 167, 130-137.
	5-Lu, T et al. Formulation and optimization of idarubicin thermosensitive liposomes provides ultrafast triggered release at mild hyperthermia and improves tumor response. J Control Release 2015, 220, 425-437
	6-Lokerse, W.J et al. In depth study on thermosensitive liposomes: Optimizing formulations for tumor specific therapy and in vitro to in vivo relations.
	Biomaterials 2016, 82, 138-150. 7-Li, L et al. Mild hyperthermia triggered doxorubicin release from optimized stealth thermosensitive liposomes improves intratumoral drug delivery and
	efficacy. J Control Release 2013, 168, 142-150. 8-Li, L et al Triggered content release from optimized stealth thermosensitive liposomes using mild hyperthermia. J Control Release 2010, 143, 274-279.
Project Title:	Evaluation of immune stimulatory effect of heat and chemotherapy in hyperthermia
	triggered drug delivery
Abstract:	Liposomes have shown great capability in formulation, reduction of side effects and enhancing pharmacokinetics
	of chemotherapeutics by stable encapsulation of chemotherapeutics and long circulating properties. However,
	effective drug delivery at the cellular level by means of such preparations is still unsatisfactory (1-3). One
	promising approach is using spatiotemporal drug release by means of liposomes with the capacity for content
	release triggered by internal or external stimuli (1). Among different stimuli, interests to application of external
	heat, hyperthermia, is getting more attention and by means of advanced liposomal preparations and heating technologies high level of control over application of heat and drug release could be achieved. Mild hyperthermia
	(41-43 oC) not only can enhance drug delivery by triggering the release or increasing permeation and distribution
	of drugs into tumor interstitium (4) but also sensitizes tumor cells to the therapy. In addition to these local mild
	hyperthermia can also induce immune responses that could be used against tumor. On the other hand most of the
	commonly used cytotoxic chemotherapeutics also invade tumors by inducing immunologic cell death. In fact, this
	is under argue whether the direct toxic effect of chemotherapeutics is responsible for the antitumor effect or it is the induced immune response that eliminate cancer cells. Therefore, in treatment of tumor by temperature
	sensitive liposomes (TSL), there are two different stimuli that stimulate immune response by different pathways
	and importantly different timings.
	While in our previous studies we enhanced the antitumor activity of TSL+ hyperthermia by optimizing liposomal
	preparations or heat application (5-8) in this project we want to evaluate how immune system could be harnessed
	in favor of tumor regression and not tumor growth and progression. We argue that immune responses induces by each arm may interfere with each other and therefore, their
	combination may not necessarily be synergistic or even additive. For example while immunogenic cell death
	mediated by therapeutic agents is in favor of anti-tumor immune response, suppression of immune system
	followed by administration of high dose of chemotherapeutics may results in opposite responses favoring tumor
	growth. Therefore, knowing the pathways, mediators and timing of immune responses provoked by these stimuli
	and when combined with each other enable proper control over treatments of tumor. Additionally, knowing these
	pathways suggests what kind of immunomodulatory agents can boost the overall therapeutic effect and to achieve such impact when is best to prescribe.
	In this project we want to evaluate the local and systemic immune reactions followed by treating mouse model of
	melanoma tumor by either local mild hyperthermia alone or TSL containing doxorubicin or idarubicin plus local
	application of heat. And later improve the therapeutic activity by adjusting drug dose, dose schedule, duration of
	hyperthermia and finally using immune modulators.
	This could be done in two in vitro and in vivo settings using protein analysis techniques such as SDS-PAGE, western blotting and proteomic analysis. immunohistochemistry analysis of treated tumors, confocal microscopy and
	intravital imaging.
Requirements of	• We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team
candidate:	work to tackle large scientific questions and thus requires a student with good communication skills.
	 Master degree or MD Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific
	part of your scholarship proposal)
	 English language requirement: English speaking countries & Netherlands: no requirement
	 Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Pathology Erasmus MC, and Radiotherapy, Amsterdam UMC
Supervisor	Associate Professor, head LEO, head NICE, Timo L.M. ten Hagen
information:	Email: <u>t.l.m.tenhagen@erasmusmc.nl</u>
	Assistant professor dr. Arlene L. Oei
world no 32 Oncology	Email: <u>a.l.oei@amsterdamumc.nl</u>
	Selected publications:
	- J Nanobiotechnology, Doi: 10.1186/s12951-021-00846-z
	 Cancers, 2020. Doi: 10.3390/cancers12030582. Biol Proced Online, Doi: 10.1186/s12575-019-0114-0
	- Advanced drug delivery reviews, 2019. Doi: 10.1016/j.addr.2020.01.003
	- Int J Nanomedicine, Doi: 10.2147/IJN.S190736
	 Int J Mol Scie, 2018. Doi: 10.3390/ijms19082420 Radiation Oncology, 2017. Doi: 10.1186/s13014-017-0813-0
	- Cancer Research, 2015. Doi: 10.1158/0008-5472.CAN-15-0816
Project Title:	Exploring the role of HPV in treatment response for cervical cancer
Abstract:	HPV is a common sexually transmitted virus that can lead to different types of cancer, including
	cervical cancer. In fact, more than 95% of cervical cancers are HPV-positive. To reduce cervical
	cancer incidence, HPV vaccines have been developed which are estimated to prevent 70-85% of
	cervical cancer. However, according to the World Health Organization, vaccination will only deliver
	a 0.1% reduction in cervical cancer mortality up to 2030 (WHO, 2021). At present, the 5-year
	overall survival of patients with localized cervical cancer is approximately 92%. Unfortunately, this
	percentage rapidly drops to 56% for patients with regional disease and to only 17% for patients with distant (metastasized). Thus, we are not yet close to eliminating the burden that cervical
	cancer imposes on women worldwide. In fact, there is clear need to develop novel treatment
	strategies for patients, particularly those with non-localized cervical cancer.
	The development of novel therapies depends on a better understanding of the disease. We
	hypothesize that the HPV viral load in cervical cancer determines immune responsiveness to anti-
	cancer treatments. More insights on the meaning of HPV viral load can be decisive for choice of
	treatment. To that end tumor (immuno)biology to radiotherapy, chemotherapy, hyperthermia and
	immune modulators needs to be thoroughly investigated in both in vitro an in vivo models in
	response to improve treatment strategies.
	3D-beads Organoids Cervical cancer biopsies
	Seal () States and the seal of the seal o
	ki67 p16
	Kié7 CO3
	PD-1 PD-1 PD-1 PD-1 PD-1
	Figure: Cervical cancer cell lines will be used in 3D-cultures; patient derived organoids are made for cervical
	tumor biopsies to study treatment responses in vitro; patient material is also used for quantification of
	immune cells to be correlated to treatment outcome.
Requirements of	 We are looking for a highly motivated, hardworking student, who has completed a BSc and MSc in biomedical sciences or a related studies, to join our team. In vitro and in vivo experiences are a pre.
candidate:	• A good command of English is required. English speaking countries & Netherlands: no requirement; other countries: IELTS 7.0
	 (min. 60.0 for all subs) or TOEFL 100 (min. 20 for all subs). We offer: supervision, lab facilities and cover laboratory costs.
	 The scholarship will have to cover: your salary and living expenses.

School/Department:	Department of Pathology Erasmus MC, and Radiotherapy, Amsterdam UMC
Supervisor	Associate Professor, head LEO, head NICE, Timo L.M. ten Hagen
information:	Email: <u>t.l.m.tenhagen@erasmusmc.nl</u>
	Assistant professor dr. Arlene L. Oei
world no 32 Oncology	Email: <u>a.l.oei@amsterdamumc.nl</u>
	Selected publications:
	- Cancers, 2020. Doi: 10.3390/cancers12030582.
	 Adv Drug Deliv Rev. Doi: 10.1016/j.addr.2020.03.006
	- Advanced drug delivery reviews, 2019. Doi: 10.1016/j.addr.2020.01.003
	- Int J Nanomedicine. Doi: 10.2147/IJN.S96123
	- Int. J. of Hyperthermia, 2019. Doi: 10.1080/02656736.2019.1685686
Project Title:	Studying the abscopal effect of thermoradiation in a triple negative breast cancer
	mouse model
Abstract:	Surgery, radiotherapy, and chemotherapy can successfully achieve control of primary breast tumours. However, many patients progress with disease recurrence and metastasis, which are refractory to treatment and correlated with (very) poor prognosis. Triple negative breast cancers, representing about 15-20% of all breast cancers, recur more rapidly (2.6 vs. 5.0 years) and are associated with lower overall survival than other breast cancers (4.2 vs. 6 years). About 10-15% of all breast cancer patients suffer from an aggressive form and will develop metastases within 3 years after diagnosis of the primary tumour. While radiotherapy and hyperthermia have been successful to treat breast cancer recurrence, a new strategy to target metastases is needed. The role of the immune system in tumor progression and response to therapy has received considerable attention. Recruitment of sufficient T-cells remains a challenge in immunologically cold tumours, such as in most triple negative breast cancers. Evidence suggests focal radiotherapy and hyperthermia can induce an abscopal effect. We aim to better understand the abscopal effect to determine e.g. the cytokine release that triggers the immune response after different radiation schedules and hyperthermia doses; and subsequently effects on cell migration, colony formation and viability.
	Cytokine release after treatment cellsCytokine release after treatment primary tumorPrim. tumor - CD3Treatment of tumor cellsChanges in cell charactersisticsImmune response & Effect on distant tumorImmune response & of tumorFigure: In vitro experiments will be used to study changes in cell characteristics after various treatment combinations and treatment schedules, in particular cytokine release and immune related cell surface receptors. In animal models the abscopal effect will be studied by treatment of the primary tumor and measuring tumor growth of the distant tumor. Subsequently mechanisms of action will be elucidated to outling to the primary tumor and measuring tumor growth of the distant tumor. Subsequently mechanisms of action will be elucidated to outling to the tumor tumor.
Requirements of	 explain treatment responses. We are looking for a highly motivated, hardworking student, who has completed a BSc and MSc in biomedical sciences or a
candidate:	 related studies, to join our team. In vitro and in vivo experiences are a pre. A good command of English is required. English speaking countries & Netherlands: no requirement; other countries: IELTS 7.0 (min. 60.0 for all subs) or TOEFL 100 (min. 20 for all subs). We offer: supervision, lab facilities and cover laboratory costs.
	The scholarship will have to cover: your salary and living expenses.

School/Department:	Department of Pathology Erasmus MC
Supervisor information:	Prof dr Adriaan B. Houtsmuller, <u>a.houtsmuller@erasmusmc.nl</u>
	Assoc. Prof dr Timo L.M. ten Hagen , <u>t.l.m.tenhagen@erasmusmc.nl</u>
world no 32 Oncology	Dr. Ann L.B. Seynhaeve, <u>a.seynhaeve@erasmusmc.nl</u>
	Website: <u>www.erasmusmc.nl</u> , <u>www.molmed.nl</u>
	Grants: Mrace Most important publications regarding this program.
	Most important publications regarding this program: 1)Biol Proced Online. 2020 Feb 1;22:3. doi: 10.1186/s12575-019-0114-0
	2)Sci Rep. 2018 Jun 25;8(1):9596. doi: 10.1038/s41598-018-27943-8.
	3)J Vis Exp. 2018 Jan 19;(131):55115. doi: 10.3791/55115. 4)Cancer Res. 2007 Oct 1;67(19):9455-62. doi: 10.1158/0008-5472.CAN-07-1599.
Project Title:	Investigating synchronization and impact of pericyte interacting with endothelial
	cells during angiogenesis.
Abstract:	Pericytes have long been neglected in research and were even believed to be absent in the
	tumor-associated vasculature. These cells are closely associated with endothelial cells and are
	important to form a functional blood conducting network in normal as well as in tumor
	development. While presence of pericytes has been documented in the past, and is reviewed
	by Simms in 1986, focused investigation into these cells is more recent as well as therapeutic
	recognition. Tumors need vessels to grow and, as we observed that tumor-associated pericytes
	are differently expressed in various tumor types, the presence or absences of pericytes can
	have implications for tumor development and therapy. We recently observed that pericyte
	motion, along different vascular tubes (i.e. growing, newly formed and established), proceeds
	via a clear synchronized pattern. At the position of an emerging endothelial sprout, the nearby
	pericytes are moving away along the existing tube to later re-emerge when the endothelial
	sprout moves further into the tissue. Also, pericytes form a front at a specified distance from
	the migrating endothelial tip cell implying a strong forward-driving synchronized
	communication between pericytes and adjacent endothelial stalk cells. Next to that, velocity
	seemed to be determined by a pericyte – endothelial cell synchronized interacting signal. Many
	questions are still not completely answered and proven.
	where do angiogenic pencytes orginate from? what
	determines interaction of pericytes with endothelial cells
	and what molecular and/or biological pathways drives
	these cells? How important is this interaction in the
	establishment of a functional vasculature and in successful
	anti-cancer therapy. What are the consequences when
	this interaction is lost? We want to explore the biological
	implications of pericyte - endothelial cell interaction in
	more detail and investigate the consequences when
	communication between pericytes and endothelial cells is
	lost. As pro- as well as anti-vascular processes are
	important in cancer treatment a better understanding of
	the close relationship between pericytes and endothelial
	cells is of critical value.
	cells is of critical value. Schematic overview of the research direction. We want to investigate the biological behavior and genetic signaling of pericytes interacting
	Schematic overview of the research direction. We want to investigate
	with endothelial cells in angiogenesis and tumor therapy.
Requirements of	 We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team work to tackle large scientific questions and thus requires a student with good communication skills. As
candidate:	mice models are a major part of the experimental set-up affinity to work with animals is required.
	Master degree or MD
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship propaga))
	 scientific part of your scholarship proposal) English language requirement:
	English speaking countries & Netherlands: no requirement
	Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Pathology Erasmus MC
Supervisor	Prof dr Adriaan B. Houtsmuller, <u>a.houtsmuller@erasmusmc.nl</u>
information:	Assoc. Prof dr Timo L.M. ten Hagen , <u>t.l.m.tenhagen@erasmusmc.nl</u>
	Dr. Ann L.B. Seynhaeve, <u>a.seynhaeve@erasmusmc.nl</u>
world no 32 Oncology	Website: <u>www.erasmusmc.nl</u> , <u>www.molmed.nl</u>
world no 32 Oncology	Grants: Mrace
	Most important publications regarding this program:
	1)Seynhaeve ALB, ten Hagen TL, Theranostics. 2020
	2)Seynhaeve ALB, ten Hagen TL. Sci Rep. 2018
	3)ten Hagen TL, Oncotarget. 2016
	4)ten Hagen TL, Nat. Protoc. 2015
	5)Seynhaeve AL, ten Hagen TL, J. Controlled Release. 2013 6)Seynhaeve AL, ten Hagen TL, Cancer res. 2008
	7)Houtsmuller AB. Sci Rep. 2019
	8)Houtsmuller AB, Nat Commun. 2016
	9)Houtsmuller AB, Sci Rep. 2015
Project Title:	Investigation the association between endothelial cells and mural cells in
	angiogenesis
Abstract:	Angiogenesis, the formation of new blood vessels, is essential for the proper development of
	tissues. Endothelial cells form the inner lining providing a dynamic barrier between underlying tissue
	and blood. Vascular mural cells are wrapped around the endothelial tube and are considered as
	stabilizing cells: control contractility and regulate endothelial proliferation. Vascular mural cells can
	be subdivided in vascular smooth muscle cells (vSMC), surrounding the larger vessels, and pericytes
	in smaller capillaries although some vessels have mural cells with properties between vSMC and
	pericytes. This distinction is more difficult in the tumor as typical properties separating arteries and
	veins are lost due to the more rapid and chaotic vessel growth. The study of angiogenesis is
	predominantly focused on endothelial cells and much less is known of mural cells. However, mural
	cells play a fundamental role in normal as well as pathological angiogenesis and are crucial for
	endothelial survival. The complex molecular association between both cells suggests that pericytes
	are more than just supporting cells. Functionality, ontogeny and identity are not fully understood
	and as there is no single common marker available to define vSMC and pericytes this makes it a
	more challenging cell type to investigate. We argue that mural cells are equally important to
	establish a functional vascular network and the cellular and molecular interaction between these
	cells will be studied. To do this we developed intravital microscopy using transgenic mice in which
	we can follow the dynamic nature of these cells in a 4D (XYZ+T, time dimension) manner. Also 2D
	and 3D in vitro cell cultures and ex vivo material will be used to study all steps in angiogenesis.
	Sprouting endothelial cells and pericytes
	a Figure: High resolution 4D
	intravital imaging of sprouting
	endothelial cells and pericytes.
	(a) Shown are 70 μm
	subsequential maximal
	projections of endothelial cells
	T = 0 hr $T = 12 hrs$ $T = 24 hrs$ $(eNOStagGFP in green) and$ aii
	pericytes (Cspg4-DsRed in red)
	in a B16BL6 melanoma tumor.
	T = 0 hr T = 24 hrs T = 24 hrs T = 24 hrs T = 24 hrs (ai, aii) Zoom-in showing Endothelial cells x Pericytes
	spatial and temporal dynamics. x represent reference points in the vasculature. Scale bar represent 100 µm.
Requirements of	• We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using team
candidate:	work to tackle large scientific questions and thus requires a student with good communication skills. As mice models are a
	 major part of the experimental set-up affinity to work with animals is required. Master degree or MD
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific
	part of your scholarship proposal)
	English language requirement:
	English speaking countries & Netherlands: no requirement
	• Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

Department of Psychiatry

Brain disorders should be considered one of the 21st century's top global health challenges as they constitute the largest burden of disease, both within Europe and worldwide.

Our understanding of the underlying etiology and pathophysiology of mental illness is necessary to create healthy changes for future generations. Yet, the study of the human brain is often challenging and difficult due to high complexity of this organ and the multifactorial nature of emotions and cognition.

Furthermore, the stigma of mental illness remains a profoundly significant barrier to early-intervention and treatment continuity, thereby perpetuating the consequences of psychiatric illness for patients, families, healthcare providers, and society.

Therefore, to address these complementary and interconnected aspects of mental illness, our department has undertaken specific areas of intense research focus within our research program from 'bench to bedside to society'.

Mission statement

Our mission is to innovate and optimise the diagnosis, treatment and prevention of severe mental health disorders in a medical context. The research conducted herein comprises applied, clinical and translational studies.

The research of the Department of Psychiatry focusses on:

- Neurobiology of Mood & Psychotic Disorders;
- Applied social and forensic psychiatry;
- Medical psychology.

Our scientific research is organized into three main research lines that, each with their specific area/ focus of interest, are distinguished by their complementary methodological approaches. The three research lines cooperate naturally.

Six examples illustrating the research carried out at the department:

- Bouwkamp CG, Kievit AJA, Markx S, Friedman JI, Zutven L van, Minkelen R van, Vrijenhoek T, Xu B, Sterrenburg-van de Nieuwegiessen I, Veltman JA, Bonifati V, Kushner SA. Copy number variation in syndromic forms of psychiatric illness: the emerging value of clinical genetic testing in psychiatry. *Am J Psychiatry 2017; 174: 1036-1050.*
- Grootendorst-van Mil, N. H., Bouter, D. C., Hoogendijk, W. J. G., van Jaarsveld, S. F. L. M., Tiemeier, H., Mulder, C. L., & Roza, S. J. The iBerry study: a longitudinal cohort study of adolescents at high risk of psychopathology. *European Journal of Epidemiology*, 2021; 36(4), 453–464.
- Influence of age on ECT efficacy in depression and the mediating role of psychomotor retardation and psychotic features. Heijnen WTCJ, Kamperman AM, Tjokrodipo LD, Hoogendijk WJG, van den Broek WW, Birkenhager TK. J Psychiatr Res. 2019 Feb;109:41-47. doi: 10.1016/j.jpsychires.2018.11.014. Epub 2018 Nov 15.
- 4. Sharma V, Bergink V, Berk M, Chandra PS, Munk-Olsen T, Viguera AC, Yatham LN. Childbirth and prevention of bipolar disorder: an opportunity for change. *Lancet Psychiatry 2019; 6(9): 786-792.*
- 5. Vrij FM de, Bouwkamp CG, Gunhanlar N, Shpak G, Lendemeijer B, Baghdadi M, Gopalakrishna S, Ghazvini M, Li TM, Quadri M, Olgiati S, Breedveld GJ, Coesmans M, Mientjes E, Wit T de, Verheijen FW, Beverloo HB, Cohen D, Kok RM, Bakker PR, Nijburg A, Spijker AT, Hassmans PMJ, Hoencamp E, Bergink V, GROUP Study Consortium, Vorstman JA, Wu T, Olde Loohuis LM, Amin N, Langen CD, Hofman A, Hoogendijk WJ, Duijn CM van, Ikram MA, Vernooij MW, Tiemeier H, Uitterlinden AG, Elgersma Y, Distel B, Gribnau J, White T, Bonifati V, Kushner SA. Candidate GSPG4 mutations and induced pluripotent stem cell modeling implicate oligodendrocyte progenitor cell dysfunction in familial schizophrenia. *Mol Psychiatry 2019; 24(5): 757-771.*
- 6. Wierdsma AI, Mulder CL. Cost sharing does not lead to an overall increase of involuntary commitments in the Netherlands. *JAMA Psychiatry 2018; 75(2): 213.*

Department of Psychiatry

School/Department:	Department of Psychiatry Erasmus MC
Supervisor information:	Nina Grootendorst, MD PhD, psychiatrist
	Email: <u>n.grootendorst@erasmusmc.nl</u>
world no 28 in Social Sciences	Website: <u>psych.nl</u> ; <u>iberrystudy.nl</u>
& Public Health	• Grants:
world no 58 in	- >1M euro of national funding for the cohort infrastructure and PhD projects
Psychiatry/Psychology	Most important publications:
	- Eur J Epidemiol. 2021
	 Psychiatry Res. 2018 BMJ Open. 2017
	- Front Psychiatry. 2018
	- J Pediatr. 2015
	- J Psychiatr Res. 2014
Project Title:	The Z factor: Adolescent Mental Health in Contemporary Society
Abstract:	Over the last decades there has been a modest but marked increase of especially common mental health problems of depression and anxiety (Mojtabai et al 2016). In particular adolescents are vulnerable for mental health problems as three-quarters of common mental health problems emerge before the age of 25 years old. At this moment, Generation Z, those born within the past 20 years, is about to enter adulthood. My research group studies the influence of common societal factors on the development of this generation. Although mental health is often considered a personal matter, mental health is affected by a combination of biological, psychological, and societal factors. The heavy influence of society in this intersectionality is often underexposed. Specific topics taken along include the influence of urbanicity on development of psychotic symptoms and drug use, the effects use of social media on sleep, the potential bidirectional relationship of financial debts and psychopathology and climate anxiety. Also, given the cross-diagnostic design, we are interested to study the phenomenology of suicidal behavior, the development of personality disorders and the prodromal phase of psychotic disorders. This project is imbedded in the iBerry cohort, a cohort of 1,022 adolescents at high risk for psychopathology in the greater Rotterdam area, the Netherlands (Grootendorst et al 2021 Eur J Epid). This cohort started in 2015 in the Erasmus MC, when participants where 15 years old and will run for 10 years. Giving the complexity, explanations would require a broad biopsychosocial approach (Bolton & Gillett, 2019). To shed light on the often complex underlying mechanisms our research integrates social and epidemiological psychiatry with biological and technical techniques, for example psychomotor tasks, examination of steroid profiles in hair samples and measures of the peripheral nervous system in relation to psychopathology. In sum, the project the Z factor will likely generate targets to improve mental health of future gener
	Keywords: adolescents, population-based, psychiatry, mental health
Requirements of	 We are looking for a highly motivated, hardworking student to join our international team. Due to the nature of the project and data, strong statistical and methodological skills, good communication skills, and an interest in mental
candidate:	health are required.
	 The student should have completed an MD or MSc in Neurosciences, Psychology, Health Sciences, Epidemiology, or a related field.
	 Within the project the student will have access to the iBerry Study data, training in epidemiology and statistics, and the broader Erasmus MC research infrastructure. The scholarship will, at least, have to cover subsistence allowance and international air plane ticket. We are happy to help with the scientific part of your scholarship proposal, please contact dr. Grootendorst at <u>n.grootendorst@erasmusmc.nl</u>

The Department of Radiology & Nuclear Medicine has an extensive research network spanning the range from the development, improvement, validation, application and assessment of imaging techniques in health and various disease systems. We use state-of-the-art radiological equipment in conjunction with advanced image analysis methods that include artificial intelligence and deep learning. The department collaborates with several clinical, fundamental and epidemiological partners within Erasmus MC.

The Department of Radiology & Nuclear Medicine has the following main areas of research:

- (1) *Clinical Research*: Musculoskeletal Research Group (ADMIRE*), Neuro-, Cardiac-, Abdominal- and Lung Imaging, Nuclear Diagnosis and Therapy, Image-Guided Diagnosis and Therapy
- (2) *Fundamental and Translation Research*: Biomedical Imaging Group Rotterdam (BIGR**), Physics in CT and MR technology, Optical Molecular Imaging, Molecular Imaging and Therapy (SPECTRIM)
- (3) *Health Sciences*: Population Imaging, Pediatric Population Neuro Imaging, Assessment of Radiological Technology (ART)
- * http://www.erasmusmc.nl/admire , ** http://bigr.nl

Why choose Radiology & Nuclear Medicine?

We offer various PhD projects on advanced image technologies and/or innovative image analysis using artificial intelligence and deep learning, working with the experts in the field. Researchers of the department publish more than 300 articles in peer-reviewed journals each year, ranked with a MNCS of 2.03 (ie quality is 2x world average). Fourteen PhD students defended their thesis in 2017.

Key publications (until Oct 2018) of the department:

- A spatio-temporal reference model of the aging brain. *Neuroimage 2018:169;11-22.* See on-line demo: <u>http://agingbrain.nl</u>
- Osteoporotic Vertebral Fracture Prevalence Varies Widely Between Qualitative and Quantitative Radiological Assessment Methods: The Rotterdam Study. *J Bone Miner Res 2018:33;560-568*.
- Two-Year Outcome after Endovascular Treatment for Acute Ischemic Stroke. *NEJM 2017:376;1341-1349.*
- Change in Carotid Intraplaque Hemorrhage in Community-dwelling Subjects: A Follow-up Study Using Serial MR Imaging. *Radiology* 2017:282;526-533.
- Semiautomated registration of pre- and intraoperative CT for image-guided percutaneous liver tumor ablation interventions. *Medical Physics 2017:44;3718-3725.*

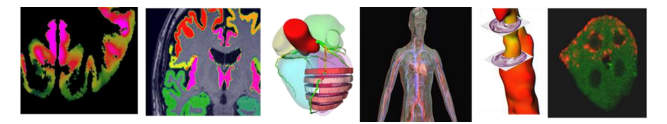
Honors & Awards (numbers from 2017):

Personal Grants/Fellowships: 12 Funded International Consortia: 11 Government Grants: 13 Grants from Charitable Organizations: 32

PPP & (Semi-)Industrial Funding: 31 Institutional Grants: 9 Travel Grants: 4

Valorization:

- Patents: https://patents.google.com/patent/WO2017010864A1/ko
- Spin-offs: Quantib BV (<u>www.quantib.com</u>)



School/Department:	Department of Radiology & Nuclear Medicine-ADMIRE, Erasmus MC
	ADMIRE-Advanced Musculoskeletal Magnetic Resonance Imaging Research Erasmus MC
Supervisor information:	Associate Professor Edwin H.G. Oei, MD, PhD: e.oei@erasmusmc.nl , www.admire-
	group.com
world no 32 Clinical Medicine	Personal Grants:
	- <i>Dutch</i> Research Council (<i>NWO</i>)
world no 36 Radiology, Nuclear	- GE Healthcare / National Basketball Association (NBA) Patellar Tendinopathy CFP 2016
Medicine & Medical Imaging	- Radiological Society of North America (RSNA) 2014
a	Most important publications:
	- Breda et al. J Magn Reson Imaging. 2020 Aug;52(2):420-430
	- De Vries et al. Semin Arthritis Rheum. 2020 Apr;50(2):177-182
	- Eijgenraam et al. Eur Radiol. 2019 Oct;29(10):5664-5672Verschueren et al. Osteoarthritis
	Cartilage. 2017 Sep;25(9):1484-1487
	Van Tiel et al., Radiology. 2016 May;279(2):523-31.
	- Van der Heijden et al. Am J Sports Med. 2016 May;44(5):1172-8
Project Title:	Analysis of advanced musculoskeletal magnetic resonance imaging (MRI) data
	from clinical and population-based studies.
Abstract:	The ADMIRE group's research focuses on imaging of common musculoskeletal diseases
	such as osteoarthritis, osteoporosis, and sports injuries, with advanced imaging
	techniques. We develop, improve, and validate innovative MRI, CT, ultrasound methods
	with the aim to identify new sensitive imaging biomarkers for pathological tissue
	processes and structural and compositional changes in tissues such as cartilage, bone,
	meniscus and tendon. We apply our novel imaging techniques in various clinical studies in
	collaboration with clinical departments. Another important research focus is on
	musculoskeletal population imaging, in which we apply MRI in the large-scale population
	based Rotterdam Study among elderly and the Generation R cohort among children and
	adolescents to study and epidemiology, genetics, and development of musculoskeletal
	diseases and body composition. The aim of this project will be to analyze existing, readily
	available, but unexplored quantitative MRI datasets acquired in clinical and population
	cohorts. The exact focus of the project and datasets to be utilized, will be defined at a
	later stage depending on the candidate's expertise and preference, but may as an
	example the assessment of bone, cartilage and meniscus quality on MRI from clinical
	osteoporosis and osteoarthritis studies, and correlation with symptoms or clinical
	outcomes. In the population imaging studies, an example would be the analysis of knee or
	hip MRI scans in the Generation R study, and correlation with risk factors and genetics.
	The project would typically entail the reading, annotation and quantitative biomarker
	extraction from acquired MRI datasets and correlating these with clinical and/or
	epidemiological data. According to the PhD student's profile and preference, the level of
	technical or analytical (MR physics, MRI analysis, deep learning) versus clinical focus will
	be defined.
Requirements of candidate:	This project requires a highly motivated, hardworking candidate with good communication skills and an affinity
4	with medical imaging and musculoskeletal disease. Given the flexibility in topic and clinical versus technical focus,
	we encourage candidates with various backgrounds including medical and technical (e.g. biomedical engineering, physics or bioinformatics) to apply.
	Master degree or MD
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the exist file part of your scholarship part could
	 the scientific part of your scholarship proposal) English language requirement:
	English speaking countries & Netherlands: no requirement
	Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Radiology & Nuclear Medicine, Erasmus MC
	BIGR-Biomedical Imaging Group Rotterdam
Supervisor information:	 Assistant Professor Dr. Esther Bron; <u>e.bron@erasmusmc.nl</u>
	Website: www.bigr.nl,https://estherbron.com/,
world no 36 Radiology, Nuclear	https://scholar.google.nl/citations?user=Mg7Q67sAAAAJ&hl=nl
Medicine & Medical Imaging	Selected publications:
	 Bron et al. Cross-Cohort Generalizability of Deep and Conventional Machine Learning for MRI- based Diagnosis and Prediction of Alzheimer's Disease, NeuroImage: Clinical, 2021 <u>https://doi.org/10.1016/j.nicl.2021.102712</u> Li et al. Longitudinal diffusion MRI analysis using Segis-Net: a single-step deep-learning framework for simultaneous segmentation and registration, <i>NeuroImage</i>, 2021 <u>https://doi.org/10.1016/j.neuroimage.2021.118004</u>
	 Venkatraghavan et al. Disease Progression Timeline Estimation for Alzheimer's Disease using Discriminative Event Based Modeling, <i>NeuroImage</i>, 2019. <u>https://arxiv.org/abs/1808.03604</u> Bron et al. Standardized evaluation of algorithms for computer-aided diagnosis of dementia based on structural MRI: the CADDementia challenge. <i>NeuroImage</i>, 2015. <u>https://caddementia.grand- challenge.org/</u>
Project Title:	Neuroimage Analysis and Machine Learning
Abstract:	Brain diseases – including dementia and stroke – impose an enormous burden to the individual and to society. As a consequence, there is an urgent need to develop effective preventive and therapeutic strategies. It is therefore essential to improve the understanding of the progression of diseases, patient selection in clinical trials, and patient monitoring in clinical practice and clinical trials. Neuroimage analysis and machine learning play a herein a crucial role, i.e. for developing robust quantitative brain imaging biomarkers and for developing data-driven models for diagnosis and prediction. PhD projects on the following topics are offered: <i>Predictive modeling of Alzheimer's disease</i> – In our research, we develop innovate diagnostic and prediction models using spatiotemporal modeling and state-of-the-art machine learning and deep learning approaches. For this we analyze of thousands of brain MRI scans and clinical data from several large clinical practice as this is hampered by the integration of multimodal biomarkers, heterogeneity of the disease and differences between datasets. In this project, we aim develop methods that can be translated towards clinical practice focusing on novel technology, multidisciplinary collaboration, objective performance evaluation beyond accuracy. <i>The baby brain pipeline: MRI analysis in craniosynostosis</i> – Syndromic craniosynostosis is a congenital disorder in which several skull sutures close prematurely, causing skull and facial anomalies. The Dutch Craniofacial Center at the Erasmus MC aims to get a better understanding of the disease process and its consequences, particularly relating to visual,
	behavioral and neurocognitive functioning. It is yet unclear whether surgery of these children is beneficial. We hypothesize that in some patients refraining from surgery might result in similar outcome, but this cannot yet be proven. We aim to use advanced MRI techniques to study the impact of craniosynostosis on the structure and function of the brain. For the analysis of these brain scans, in small children with brain deformations, no automated approaches exist. The proposed project aims at development of dedicated image analysis tools for children with craniosynostosis.
Requirements of candidate:	 This project requires a highly motivated, hardworking candidate with good communication skills, who likes to become part of our international team. Master degree in a technical discipline preferably with an affinity for medical applications (medical physics, biomedical engineering, physics, computer science, engineering,) Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) English language requirement: English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Radiology & Nuclear Medicine, Erasmus MC
	BIGR-Biomedical Imaging Group Rotterdam
Supervisor information:	Prof dr Wiro Niessen,: <u>w.niessen@erasmusmc.nl</u> <u>www.bigr.nl</u>
	Personal Grants:
world no 36 Radiology,	Wiro Niessen is (co-PI) of numerous Dutch and European research grants, including on
Nuclear Medicine & Medical	Imaging Genetics (1 MEuro), Radiomics (600 kEuro). He received personal VICI
Imaging	
	grants (1.25 MEuro) and Simon Stevin award (500 kEuro). Total research funding
	over last 10 years is more than 15 MEuro. He has supervised 42 PhD students.
	Most important publications:
	 Hofer, E.et al 2020. Genetic correlations and genome-wide associations of cortical structure in general population samples of 22,824 adults. Nature Communications, 11(1), pp.1-16.
	 Van der Lee SJ et al. Gray matter heritability in family-based and population-based studies using voxel-based morphometry.
	Human Brain Mapping. 2017;38(5):2408-23.
	 Wang, J. et al2019. Gray matter age prediction as a biomarker for risk of dementia. Proceedings of the National Academy of Sciences, 116(42), pp.21213-21218.
	- Hibar DP et al. Novel genetic loci associated with hippocampal volume. Nature Communications. 2017;8.
	 Roshchupkin GV et al. Heritability of the shape of subcortical brain structures in the general population. Nature Communications. 2016;7.
	- Santos EMM et al. Observer variability of absolute and relative thrombus density measurements in patients with acute
	 ischemic stroke. Neuroradiology. 2016;58(2):133-9. Roshchupkin GV et al. HASE: Framework for efficient high-dimensional association analyses. Scientific Reports. 2016;6.
	 Roshchupkin GV et al. Fine-mapping the effects of Alzheimer's disease risk loci on brain morphology. Neurobiology of Aging.
	2016;48:204-11.
	 Niessen WJ. MR brain image analysis in dementia: From quantitative imaging biomarkers to ageing brain models and imaging genetics. Medical Image Analysis. 2016;33:107-13.
	 Huizinga W et al. PCA-based groupwise image registration for quantitative MRI. Medical Image Analysis. 2016;29:65-78.
Project Title:	Distributed Machine Learning in application for large-scale omics studies
Abstract	Artificial Intelligence field has seen dramatic advances in the past few years with much
	excitement around the use of deep learning (DL), many-layered convolutional neural networks
	(CNN). The world has witnessed striking advances in the ability of machines to understand and
	manipulate data, including images, language, and speech. CNN showed ability to detect a
	complex pattern in high-dimensional data, but also are able to integrate data from various
	resources by having many input channels into neural network. Human genetics can benefit
	immensely from DL. However, the application of AI in genetics analysis is still quite limited.
	The main issue is the restriction for data sharing between cohorts and loss of power, compare
	to the pooled analysis.
	Distributed Learning is a distributed machine learning approach which enables model training
	on a large corpus of decentralized data.
	The main goal of this project is to develop new distributed learning framework for multi-
	center genetics analysis in collaboration with NVIDIA company, which will be able to utilize
	machine learning approaches and increase power of gene discovery. We aim to apply these
	methods on large datasets from population-based Rotterdam study, UK Biobank as well as
	within world-wide genetics consortiums.
Requirements of	We are looking for a highly motivated, hardworking student to join our very international team. Successful candidates are expected to have a strong quantitative or computer science background, excel at critical thinking, with a strong motivation to
candidate:	engage in the development and application of advanced analytical methods.
	• Master degree in mathematics, computer science, statistics, bioinformatics, physics, electrical engineering, or in an equivalent
	discipline. •Strong knowledge of: Python.
	•Experience with machine learning and deep learning methods.
	•Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal)
	English language requirement:
	English speaking countries & Netherlands: no requirement
	Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Radiology & Nuclear Medicine, Erasmus MC
	BIGR-Biomedical Imaging Group Rotterdam
Supervisor information:	Associate Professor Dr. ir. Stefan Klein; <u>s.klein@erasmusmc.nl</u>
	Website: https://scholar.google.nl/citations?user=iaAFK0MAAAAJ
world no 36 Radiology, Nuclear	• Selected publications:
Medicine & Medical Imaging	- Venkatraghavan et al. Disease Progression Timeline Estimation for Alzheimer's Disease using
	Discriminative Event Based Modeling, NeuroImage, 2019. https://arxiv.org/abs/1808.03604
	- Sun, Niessen, Klein. Randomly perturbed B-splines for nonrigid image registration. IEEE
	Transactions on Pattern Analysis and Machine Intelligence, 2017. <u>CSC funded</u>
	- Huizinga et al. PCA-based groupwise image registration for quantitative MRI. <i>Medical Image</i>
	Analysis, 2016.
	- Bron et al. Standardized evaluation of algorithms for computer-aided diagnosis of dementia based on structural MRI: the CADDementia challenge. <i>NeuroImage</i> , 2015.
	https://caddementia.grand-challenge.org/
	 Klein, Staring et al. Elastix: a toolbox for intensity-based medical image registration. <i>IEEE</i>
	Transactions on Medical Imaging, 2010. (>2500x cited, software used by researchers and
	companies worldwide, <u>www.elastix.isi.uu.nl</u>)
Project Title:	Image Analysis and Machine Learning
Abstract:	We develop advanced image analysis methods and machine learning approaches to
	extract more information from medical images than can be seen by the naked eye. PhD
	projects on the following topics are offered:
	<u>Radiomics for precision cancer medicine</u> - Radiomics is a big-data analytics technique, in
	which hundreds of candidate features are calculated from imaging data and annotated
	tumour contours, quantifying location, shape and appearance of the tumour. Using
	machine-learning algorithms, such as SVMs or deep neural networks, these computational
	features are combined into predictive models, also called 'radiomics signatures'. At
	Erasmus MC, we have access to unique datasets that allow development of novel
	radiomics signatures that could aid the diagnosis and treatment of cancer.
	Disease progression modelling of neurodegenerative diseases – Alzheimer's Disease and
	related disorders of the brain are a major challenge in the ageing population worldwide.
	Development of novel curative treatments is hampered by the heterogeneity of the
	disease, lack of reliable tools for early and differential diagnosis, and limited insight in the
	various disease progression patterns. In our research, we develop innovate computer-
	aided diagnosis methods and data-driven disease progression models, using
	spatiotemporal analysis of thousands of brain MRI scans.
	Image analysis and machine learning for osteoarthritis – Osteoarthritis is the most
	common degenerative disorder of the knee joint. Reliable methods for early diagnosis,
	fine-grained disease staging, and accurate patient stratification are urgently needed to
	improve patient care. MRI provides 3D visualization of multiple tissues in and around the
	knee joint, and holds great promise as a basis for detailed phenotyping and spatial
	mapping of pathology. In collaboration with the ADMIRE group (headed by Dr. Oei), we
	develop methods for quantitative MRI analysis, and study the relation of MRI markers
Deminenceste ef constitutes	 with clinical, biochemical, and genetic markers. This project requires a highly motivated, hardworking candidate with good communication skills, who likes to become
Requirements of candidate:	part of our international team.
	• Master degree in a technical discipline (physics, mathematics, computer science, engineering, etc.)
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal)
	 scientific part of your scholarship proposal) English language requirement:
	• English speaking countries & Netherlands: no requirement
	• Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Radiology & Nuclear Medicine, Erasmus MC BIGR-Biomedical Imaging Group Rotterdam
Supervisor information:	Dr. Theo van Walsum
	Email: t.vanwalsum@erasmusmc.nl
World no 23 in Cardiac &	Website: www.bigr.nl , www.bigr.nl/people/TheovanWalsum
Cardiovascular Systems	Most important publications:
world no 36 Radiology, Nuclear Medicine & Medical	- autoTICI: Automatic Brain Tissue Reperfusion Scoring on 2D DSA Images of Acute Ischemic Stroke Patients, IEEE TMI 2021
Imaging	 Automatic collateral scoring from 3D CTA images, IEEE TMI 2020 Automated quantification of bileaflet mechanical heart valve leaflet angles in CT images, IEEE TMI 2018
	 Quantitative analysis of geometry and lateral symmetry of proximal middle cerebral arteryJSCD 26(10), 2017
	- Automatic segmentation and quantification of the cardiac structures from non-contrast- enhanced cardiac CT scans, PMB 62(9), 2017
	 Classification of hemodynamically significant stenoses from dynamic CT perfusion and CTA myocardial territories MP 44(4), 2017
	- Epicardial fat volume and the risk of atrial fibrillation in the general population free of cardiovascular disease, JACC: Cardiovascular imaging, 2017
Project Title:	Quantitative Imaging Biomarkers for Cardiovascular Diseases
Abstract:	Cardiovascular disease is one of the major health problems in the western world. Whereas
	treatment options are growing, there is still much unknown on diseases and optimal
	treatment strategies. Quantitative imaging biomarkers may play an import role in this field.
	Using quantitative information from images can learn more on diseases and disease
	development, and may, based on this knowledge, also provide information for clinical decision
	making. Additionally, the large amounts of imaging data and clinical data may also be used to directly learn decision models from existing databases.
	In this research line, we are developing quantitative imaging biomarkers for cardiovascular
	diseases. We are focusing on CTA (cardiac, brain) as well as X-ray imaging modalities (the latter
	for interventional decision making), for heart disease and stroke. In this work, we are also
	more and more exploiting the power of deep learning approaches.
	Examples of recent studies from our group in this field are listed above.
Requirements of	• We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using
candidate:	 team work to tackle large scientific questions and thus requires a student with good communication skills. Master degree in an engineering discipline
	 Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal)
	English language requirement:
	 English speaking countries & Netherlands: no requirement Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)

School/Department:	Department of Radiology & Nuclear Medicine, Erasmus MC				
	BIGR-Biomedical Imaging Group Rotterdam				
Supervisor information:	Dr. Theo van Walsum				
	Email: t.vanwalsum@erasmusmc.nl				
World no 23 in Cardiac &	Website: www.bigr.nl , www.bigr.nl/people/TheovanWalsum				
Cardiovascular Systems	Most important publications:				
world no 36 Radiology, Nuclear Medicine & Medical	- Virtual extensions improve perception-based instrument alignment using optical see-through devices. IEEE TVCG, 2021				
Imaging	 Dynamic coronary roadmapping via catheter tip tracking in X-ray fluoroscopy with deep learning based Bayesian filtering, MedIA 61, 2020 				
	- Ultrasound aided vertebral level localization for lumbar surgery, IEEE TMI 36(10)				
	- A Hidden Markov Model for 3D Catheter Tip Tracking With 2D X-ray Catheterization Sequence and 3D Rotational Angiography, IEEE TMI 36(3)				
	 Non-rigid registration of liver CT images for CT-guided ablation of liver tumors, Plos One 11(9) 4D Ultrasound tracking of liver and its verification for tips guidance, IEEE TMI 35(1) 				
	- Automatic online layer separation for vessel enhancement in X-ray angiograms for				
	percutaneous coronary interventions, MedIA 39				
Project Title:	Trackerless navigation approaches for interventional radiology and cardiology				
Abstract:	Minimally invasive interventions are good for patient and society. Compared to conventional				
	surgery, minimally invasive interventions give reduced trauma, leading to benefits for patient				
	and society. These advantages come at the expense of the physician, who often lacks direct				
	eyesight and tactile feedback during the interventions.				
	Surgical navigation systems, which link the patient to pre-operative imaging information, and				
	which are equipped with systems to track instrument and patient motion, have been utilized				
	in e.g. neuro, spine and orthopedics surgery to support the physician in minimally invasive				
	interventions.				
	Purpose of the research in this project is to develop technology that permits navigation				
	approaches in soft tissue interventions, such as percutaneous coronary interventions and liver				
	interventions (tumor ablations). To this end, we are utilizing imaging information (ultrasound /				
	X-ray) acquired during the procedures, and integrate pre-operative information in these				
	images. For this, advanced segmentation, registration and tracking methods have been developed, and more recently we are also exploiting deep learning methods for these				
	purposes. The publications listed above show some of the recent approaches in this line.				
	Additionally, we are investigating augmented reality approaches for navigation.				
Requirements of	We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in using				
candidate:	team work to tackle large scientific questions and thus requires a student with good communication skills.				
	 Master degree in an engineering discipline Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the scientific part of your scholarship proposal) 				
	English language requirement:				
	English speaking countries & Netherlands: no requirement				
	• Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs)				

School/Department:	Department of Radiology and Nuclear Medicine, Erasmus MC						
	Molecular Medicine						
Supervisor information:	Associate Professor Dr. Yann Seimbille, <u>v.seimbille@erasmusmc.nl</u>						
	• Website: 1) <u>https://www.erasmusmc.nl/en/research/departments/radiology-and-nuclear-medicine;</u>						
world no 32 Oncology	2) <u>https://www.erasmusmc.nl/en/research/groups/radiopharmaceutical-chemistry</u> ; 3)						
	https://www.erasmusmc.nl/en/research/researchers/seimbille-yann						
world no 36 Radiology,	• Grants:						
Nuclear Medicine & Medical	- Long-acting sstr2 antagonists and pretargeted alpha therapy, Dutch Cancer Foundation , 2019-2023						
<u>Imaging</u>	- Broad spectrum, high precision theranostic cancer therapy, Convergence kick-off grant, 2020-2022						
	- Theranostics hitting breast cancer: pointing the arrows at HER2 and GRPR, Erasmus MC Grant , 2021-2025						
	 Most important publications: Koustoulidou S, Hoorens M, Dalm S, Debets R, Mahajan S, Seimbille Y, de Jong M. <u>Cancers</u>, 2021, 13(5), 1100 						
	(https://doi.org/10.3390/cancers13051100).						
	- Chen KT, Nieuwenhuizen J, Handula M, Seimbille Y . <u>Organic and Biomolecular Chemistry</u> . 2020, 18(31), 6134-						
	6139 (<u>https://doi.org/10.1039/D0OB01222J</u>).						
	- Qiu L, Wang W, Li K, Peng Y, Lv G, Liu Q, Gao F, Seimbille Y , Xie M, Lin J. <u>Theranostics</u> . 2019, 9(23), 6962-6975						
	(<u>https://doi.org/10.7150/thno.35084</u>).						
	 Chevalier C, Stojanović O, Colin DJ, Suarez-Zamorano N, Tarallo V, Veyrat-Durebex C, Rigo D, Fabbiano S, Stevanović A, Hagemann S, Montet X, Seimbille Y, Zamboni N, Hapfelmeier S, Trajkovski M. Cell. 2015, 163, 						
	1360-1374 (https://doi.org/10.1016/j.cell.2015.11.004).						
	 Suarez-Zamorano N, Fabbiano S, Chevalier C, Stojanovic O, Colin DJ, Stevanovic A, Veyrat-Durebex C, Tarallo 						
	V, Rigo D, Germain S, Ilievska M, Montet X, Seimbille Y , Hapfelmeier S, Trajkovski M. <u>Nature Medicine</u> . 2015,						
	21, 1497-1501 (<u>https://doi.org/10.1038/nm.3994</u>).						
	- Su H, Bodenstein C, Dumont RA, Seimbille Y, Dubinett S, Phelps ME, Herschman H, Czernin J, Weber W.						
	<u>Clinical Cancer Research</u> . 2006, 12, 5659-5667 (<u>https://doi.org/10.1158/1078-0432.CCR-06-0368</u>).						
Project Title:	Theranostic agents for cancer imaging and therapy						
Abstract:	The RadioPharmaceutical Chemistry (RPC) group's research program is a molecular imaging-based						
	program focused on theranostics and multimodality imaging probes, with an emphasis on						
	developing these novel radiopharmaceuticals for clinical translation.						
	We are offering to work on a						
	project aiming at the RadioPharmaceutical Chemistry						
	development of a new						
	generation of theranostics						
	pointing at the major Achilles'						
	heels of tumors, such as the						
	fibroblast activation protein						
	alpha (FAPa) or the chemokine						
	receptor type 4 (CXCR4). The						
	new radioactive drugs will be						
	capable of providing adequate						
	diagnostic information and (Image guided surgery)						
	subsequently kill the tumor cells						
	when targeted radionuclide therapy is found appropriate. Addition of a fluorescent dye will						
	provide dual-modality imaging probes for pre-operative surgical planning and intraoperative						
	surgical guidance, whereas conjugation of a potent antineoplastic drugs will yield small-molecule						
	drug conjugates (SMDC) for targeted chemotherapy. Preclinical evaluations of our theranostics						
	will allow to identify which lead candidate could potentially be translated to the clinic.						
Requirements of	• We are looking for a highly motivated, hardworking student to join our very international team. Our strength is in						
candidate:	using team work to tackle large scientific questions and thus requires a student with good communication skills.						
	 Master degree in the field of Chemistry, Biochemistry or Pharmaceutical Sciences. Strong expertise in organic chemistry and analytical techniques (NMR, HPLC, MS) required. Experience with radiolabeling techniques and 						
	biological assays is an asset.						
	• Scholarship that will, at least, cover subsistence allowance and international air plane ticket (we could help with the						
	scientific part of your scholarship proposal)						
	English language requirement: O English speaking countries & Netherlands: no requirement						
	 Other countries: IELTS 7.0 (min 6.0 for all subs), TOEFL 100 (min 20 for all subs) 						

Department of Surgery

School/Department:	Department of Surgery, Erasmus MC				
Supervisor information:	Prof. dr. Luc van der Laan & dr. Monique Verstegen				
	I.vanderlaan@erasmusmc.nl / m.verstegen@erasmusmc.nl				
world no 13 Surgery	Selected publications:				
world no 14 Gastroenterology	- Materials Science & Engineering, 2020, Willemse, van der Laan & Verstegen, et al				
<u>& Hepatology</u>	- Transplantation, 2020, Verstegen & van der Laan, et al				
	 Cancers, 2019, van Tienderen, van der Laan & Verstegen, et al. Nature Medicine, 2017, Broutier , Verstegen, van der Laan & Huch, et al. 				
	 Nature Medicine, 2017, Broutier , verstegen, van der Laan & Huch, et al. Nature, 2016, Blokzijl, Verstegen, van der Laan & van Boxtel et al. 				
Project Title:	Exploring the regenerative potential of liver organoids in liver transplantation				
Abstract:	Although the adult liver is well-known for its regenerative capacity, the cellular events that drive this repair are pleiotropic and not fully elucidated. The two liver epithelial cell types, hepatocytes and cholangiocytes, have self-renewal capacity to maintain homeostasis and in response to liver injury. Moreover to the plasticity of epithelial cells, bipotent progenitor cells are found within the canals of Hering, the smallest branches of the biliary tree in the liver. These bipotent progenitor cells can differentiate into both mature hepatocytes and cholangiocytes. In larger bile ducts, including in the extrahepatic bile ducts, typical peribiliary glands harbor biliary progenitor cells which provide a proliferative response upon damage of the bile duct providing new cholangiocytes to restore the biliary lining. With the development of the 3D organoid culture technique, epithelial cells, including those found in the liver can be expanded <i>in vitro</i> (Huch et al, Cell, 2015) and used as model for stem cell biology and liver diseases such as Metabolic Associated Fatty Liver Disease (MAFDL) or primary liver cancer.				
	developing liver-on-a-chip technology), and liver and bile duct tissue engineering (decellulairsation techniques and extracellular matrix analysis). During liver transplantation performed in Erasmus MC, biopsies are collected from liver and extrahepatic bile duct from donor and recipient (explanted liver) to be used in research projects. These biopsies are analyzed using histological techniques (immunohistochemistry, immunofluorescence, conventional, confocal and light-sheet microscopy) and molecular biological techniques (qPCR, RNA-expression arrays and whole genome sequencing). In addition, the LGR5-positive, Wnt-responsive adult stem cells from liver and the extrahepatic bile duct, will be cultured and expanded as organoids to be used as (patient-specific) models for liver regeneration and/or disease, including primary liver cancer.				
	Main methodology and techniques: 3D biliary organoid cultures from healthy donor and				
	patient biopsies (NASH, primary liver cancer). Gene expression analysis (single cell RNA sequencing, RT-qPCR), high resolution imaging (OIC-confocal, fluorescence microscopy),				
	protein expression analysis (FACS, Immunohistochemistry, Western blotting).				
Requirements of candidate:	 We are looking for a highly motivated PhD student who has received excellent scientific and practical training in the areas of stem cell biology, transplantation medicine and/or regenerative medicine to join our research team. The student should be fluent in English (IELTS <i>min 6.0</i>), TOEFL 100 (<i>min 20 for all subs</i>). 				
	• We offer: Supervision, lab facilities and infrastructure, and training.				
	• We will cover Laboratory costs.				
	• As a candidate PhD student at Erasmus MC, your salary and living expenses will be covered by your University or Scholarship Council.				

ALASAN MEMILIH ERASMUS MC

Anda sangat disambut: kami berharap menyapa Anda sebagai mahasiswa PhD kami dan sebagai kolega kami di masa depan. Kami berharap Anda akan merasa seperti di rumah sendiri dan akan berkolaborasi dengan kami dalam langkah berikutnya dalam karier Anda.

Langkah berikutnya dalam karier Anda: gelar PhD di Erasmus MC berarti Anda memiliki 4 publikasi internasional yang dinilai rekan sejawat dan memiliki publikasi berperan penting untuk langkah berikutnya dalam karier Anda. Sebagian besar universitas mengharuskan satu publikasi atau kurang, sehingga publikasi dari Erasmus MC merupakan keunggulan signifikan (lihat hal. 3 untuk prestasi 10 PhD Indonesia terakhir).

Pendidikan & pelatihan Anda: dengan sekitar 1.500 staf ilmiah untuk kurang dari 1.250 mahasiswa PhD dan ~750 spesialis kedokteran untuk ~1.000 residen, kami memiliki rasio pembimbingan yang sangat baik. Hal yang lebih penting adalah mahasiswa PhD memiliki setidaknya 2 pembimbing di Erasmus MC dan 1 pembimbing Indonesia karena kami memilih untuk melatih Anda sedemikian rupa agar Anda dapat melanjutkan penelitian Anda di Indonesia.

Tidak perlu belajar bahasa Belanda: tidak perlu belajar bahasa Belanda – Belanda menduduki peringkat <u>pertama atas</u> <u>kefasihannya dalam bahasa Inggris</u> dalam dua tahun terakhir dan berada di peringkat 3 teratas dalam sepuluh tahun terakhir dan Rotterdam menduduki peringkat pertama di antara kota-kota di Belanda. Jadi, Anda bisa berbelanja di toko tanpa harus berbicara dalam bahasa Belanda.

Kehidupan sosial Anda: lebih dari 30% dari mahasiswa PhD kami adalah mahasiswa asing dan kami memiliki organisasi mahasiswa PhD yang aktif di <u>Erasmus MC</u> dan <u>Erasmus University Rotterdam</u>, dan kantor-kantor internasional. Tinggal di kota pelabuhan terbesar di Eropa, yang ditampilkan menduduki peringkat <u>ke-5 menurut Lonely Planet pada tahun 2016</u>, berarti Anda berjarak satu jam dari Amsterdam atau Antwerp (dengan mobil), dari Brussels (dengan kereta api), dari London (dengan pesawat terbang) atau 1,5 jam dari Berlin (pesawat terbang) atau 2 jam dari Paris (dengan kereta api).

Organisasi kami: Erasmus MC adalah salah satu dari sepuluh sekolah kedokteran terbesar di Eropa dan salah satu dari sepuluh lembaga terbesar dalam publikasi ilmu praklinis, klinis, & kesehatan yang disubsidi oleh Komisi Eropa. Kolaborasi ilmiah kami dengan mitra dari Indonesia sangat baik dan kualitas kolaborasi kami dengan ASEAN dan Indonesia (sebagaimana ditunjukkan dengan sitasi/publikasi rata-rata, lihat tabel kanan di bawah ini) sangat tinggi dibandingkan dengan universitas asing lain, yang merupakan keunggulan saat membawa kolaborasi penelitian Anda kembali ke Indonesia. Kami juga berperingkat ke-13-36 dunia untuk berbagai bidang klinis (lihat tabel di bawah ini) dan berperingkat <u>ke-30 dunia untuk ilmu biomedis</u>.

Kami melatih ilmuwan muda Indonesia dan berharap mereka akan menjadi generasi kolaborator Indonesia berikutnya. Kami berharap Anda bergabung dengan Erasmus MC dan menjadi kolega kami di masa depan di Belanda dan setelah Anda kembali ke Indonesia, karena hubungan kita tidak terputus setelah Anda memperoleh gelar Anda.

US News Ranking 2021	World rank	On the US News	Preclinical, clinical & Health Sciences 2016-2020		
Surgery	13	website,	InCites Clarivate dbase as of Oct, 5th, 2021		
Gastroenterology & Hepatology	14		University or Med School only*	publ	world impact
Public, Environmental & Occup Health	21	MC is ranked as	Erasmus MC*	24,271	2.55
Cardiac & Cardiovascular Systems	23	Erasmus	Erasmus University Rotterdam	25,746	2.52
Infectious Diseases		University	UCLA DG Med School*	15,863	2.47
	Z4 Botter	Rotterdam	Harvard University	139,589	2.37
Endocrinology & Metabolism	29	for the	Stanford University	40,396	2.32
Immunology	31	given	Johns Hopkins University	63,010	2.27
Clinical Medicine	32	subject	Johns Hopkins Medicine*	22,879	2.27
Oncology	32	rankings.	Harvard Univ Med School*	70,795	2.27
Radiology, Nuclear Med & Med Imaging	36	1000	UC San Francisco	47,712	2.22
Pharmacology & Toxicology	39		Yale University	34,241	2.21
Neuroscience & Behavior	48		UC Los Angeles (UCLA)	37,742	2.21
Cell Biology 67			University of Chicago	16,265	2.13

Erasmus MC Indonesia PhD Vacancy booklet 2021-2022 version 1, October 12, 2021;- RDO, Research Development Office, Dr Raoul Tan - Senior Advisor International Affairs