Aantal:

Requestform Genomediagnostics Nijmegen

Postal address	Patient information (*	=required field)	
Radboudumc Genome Diagnostics, route 815 Head laboratory: Dr. H. Yntema Postbus 9101 6500 HB Nijmegen The Netherlands Opening hours laboratory: 08:30 - 16:30 Tel: +31(0)24 3613799 Fax: +31(0)24 3616658 @: gen@radboudumc.nl Tel: https://order.radboudumc.nl/en/genetics	Initials [*] Last name [*] Date of birth [*] Gender [*] Ethnical background Address Postal code City Reference / MDN Is patient deceased? Date of death		
Invoice address (*=required field)			
Name [*] Address [*]		E-mail Vat number	
	N	Vachaniber	
Referring physician (*=required field Name* Hospital / Institute* Address Email* Specialism Department* Country Telephone)	Fax Copy results	
Material Material already available in our lab Date of sample collection Which kind of material will be included	DN	IA isolated from:	
Reason of the request Reason of referral Gene involved and mutation Has material of family member(s) be Is this an urgent request? Reason of urgency	en sent previously? N	ame(s)	Date of birth
Date of request:			Version: 07-2025
<u>To be filled in by the lab</u> Datum ontvangst: Paraaf ontvangst:	Ingescande pagina's:		monstersticker(s)
Materiaal: EDTA HEP	DNA Vlok Vru	uchtw. Overig:	

Family members

Relation to current patient Other familial information Will you include parental samples? Name father Name mother Parental clinical details

Which kind of parental material? Date of birth father Date of birth mother

Reason including parental samples

Informed consent

Genetic testing may reveal unsolicited findings. By requesting this test, we assume that you have discussed the possibility of such findings with the counselee (Dutch guideline for disclosure of incidental findings during genetic diagnostic testing). The material and/or data of the patient may be used if the current test does not find a cause for the clinical symptoms. In that case, a follow-up investigation related to the same clinical question can be performed later without a new request. This may occur, for example, when a new variant or gene is discovered, or if a more accurate test becomes available. The original requester will be notified about this. The laboratory will not charge for this. However, please note that there may be costs for the patient from the treating physician. The patient DOES NOT ALLOW the material and/or data to be used for additional analysis in line with the original diagostic request. Further use of the material has not (yet) been discussed.

Clinical information

Birth weight Current length Age of diagnosis Current weight Head circumference Consanguinity Expected inheritance

Clinical relevant information

Pedigree chart

Please mark the individual of this request with an arrow (\rightarrow). Designate affected family members as \blacksquare / \bullet and indicate previous sent family samples with name and date of birth.



Collaboration MUMC+

The laboratory Genomediagnostics Nijmegen closely collaborates with the Clinical Genomics Laboratory of the MUMC+ in Maastricht. A limited part of the test offer (marked 1) will be carried out by the partner laboratory in Maastricht. In that case, the diagnostic report will be issued directly by the Maastricht laboratory, but the payment will be issued by Genomediagnostics Nijmegen. Direct analyses of variants in family members detected by whole exome sequencing (WES) will be carried out by the laboratory that reported the variant originally in the index.

Material withdrawal form

Patient information/sticker

Initials	Date of birth
Last name	Gender

Material and shipping conditions

- Put name, gender and date of birth on each blood- or DNA tube. Improperly labelled samples will be refused. Excess tubes will not be stored!
- Ship samples at room temperature. Do not freeze blood samples!
- Do not submit material if patient has undergone bone marrow or stem cell transplant. Please contact lab.
- Upon cancellation the requested test will be charged completely.
- Additional information for requesting diagnostic testing at the Division of Genome Diagnostics can be found at https://www.radboudumc.nl/en/afdelingen/genetica/about-us/genomediagnostics

Service	Turnaround time	Required material	
Exome sequencing diagnostics (WES)	Exome gene panel analysis: 2-3 months Exome gene panel analysis followed by exome wide analysis (in one report): 2-3 months Rapid Trio/de novo analysis of proband and both parents: <15 business days	 Preferably: 2 x 3-6ml EDTA blood 20 ml amniotic fluid (only Rapid) or >30 ml (Rapid + QF-PCR growing cells) If there is no blood and/or amniotic fluid available: Desired amount DNA: 5ug Absolute minimum: 3ug (minimum concentration is 25 ng/ul, volume 30 ul). Prenatal material minimum is 450 ng (30 ul of 15 ng/ul) 	
Interpretation of exome data	2 months	n/a	
Array diagnostics (genome wide)	5 weeks	2 x 3-6ml EDTA blood	
Multiple gene diagnostics (Gene panels)	3-8 weeks*	2 x 3-6ml EDTA blood	
Single gene diagnostics	4-8 weeks*	2 x 3-6ml EDTA blood	
Mutation diagnostics (carrier testing)	4 weeks	Gene/array diagnostics: 2 x 3-6ml EDTA blood Chromosome diagnostics (karyotyping): 2 x 5ml Heparin blood in Natrium- or Lithium-Heparin tubes (neonates 1-2ml)	
Farmacogenetics	1-8 weeks*	2 x 3-6ml EDTA blood	
Chromosome diagnostics	2-5 weeks	Karyotyping : 2 x 5ml Heparin blood in Natrium- or Lithium- Heparin tubes (neonates 1-2ml) QF-PCR: neonates: 1-2ml EDTA blood	
FISH	2-5 weeks	2 x 5ml Heparin blood in natrium- of lithium-heparin tubes	
Urgent request	Please, contact: gen@radboudumc.nl or Tel: +310243613799, TAT depends on technique	Please, see material under requested service	
mtDNA	4-12 weeks	Cooled (none frozen) urine, we prefer 50-100ml	

* In our application system you will find the exact turnaround times for each individual test