Declaration of Informed Consent

for the morphological (morphology = structural description of blood cells), immunophenotypic (immunophenotyping = examination of the structure of cell surfaces), genetic (examination of chromosomes) or molecular (examination of individual genes) analysis and storage of the test material.

The findings obtained for you, in the opinion of the attending physician, indicate suspicion of a blood and/or bone marrow disease that may result from genetic aberration. Blood or bone marrow (or a different tissue) was taken from you in order to clarify this suspicion with a greater degree of precision. This material will be submitted to the aforementioned special practice with special laboratory for more precise analysis. Depending on the request and individual necessity, this material will be used to analyze chromosomes (carriers of genetic information), to isolate nucleic acids (constituent parts of the genome) and to test them for mutations (genetic aberrations), specifically in the cells of the blood or bone marrow. Among other things, this may lead to the discovery of congenital abnormalities that are probably without pathological significance for you, but which may be relevant to your progeny. Excess material will be stored for the purposes of verification. This material can also be of substantial benefit to research and development in the field of medical-genetic diagnostics. Accordingly, the test material may be used for these purposes as well. Your data will be used exclusively in an anonymized form for scientific testing and evaluation (possibly with scientific collaborators). The legal foundation for this is defined in Article 9 (2)(j) and in Article 89 (1) EU General Data Protection Regulation and in its Recitals 53, 156, 157 and 159, as well as in Section 27 Federal Data Protection Act (BDSG-new). Recital 33 to the EU General Data Protection Regulation also states that you are entitled to restrict your consent to particular areas of research only.

The retention period for diagnostic data is 10 years in accordance with statutory regulations. Data that is collected in an anonymized form for scientific purposes will not be erased.

Your attending physician has instructed you in the meaning and possible consequences of these tests. Naturally, all of your details and the findings of the tests are subject to medical confidentiality. Your attending physician will inform you of the findings of our tests. Any disclosure of diagnostic data to third parties (hospitals, other medical practices) requires your prior consent.

I request that the analyses recommended by my doctor for further diagnostic clarification be performed.

Last name: First name:
Place, date: Signature:

I consent to the use of excess test material for scientific purposes and to the Declaration of Informed Consent as above.

Yes [ ] No [ ]

Place, date: Signature:

You are entitled at any time, without being required to provide reasons, to object to the processing of scientific data pursuant to Article 21 EU General Data Protection Regulation. Kindly fax your objection without any formal requirements to +89 990 17 111, or send it by email to info@mll.com.

Pursuant to Section 27 (1) Federal Data Protection Act (BDSG), we are entitled, even without your consent, to collect data for scientific purposes, provided we comply with national processing regulations.