

Goals of the research network

Feature	Explanation
Number	G1
Name	Conduct research project
Description	<p>The main goal of the SFB/TRR77 project on liver cancer is to get a deeper understanding of the molecular basis of the corresponding pathogenesis. This includes aspects like</p> <ul style="list-style-type: none">• The beginning of the chronic liver disease• Progression• Final cancer stadium with metastases <p>Further, the research network aims to identify new preventive, diagnostic and therapeutic approaches.</p>
Weighting	high

Feature	Explanation
Number	G2
Name	Answer research questions
Description	<p>The research network covers, among others, the following research questions:</p> <ul style="list-style-type: none">• Which generic or specific mechanisms of chronic liver diseases and inflammatory processes predispose or initiate liver cancer?• Which molecular key events that promote or maintain can be used as tumor markers?• Which are promising targets for future therapies?
Weighting	high

Feature	Explanation
Number	G3
Name	Create, store, and retrieve data
Description	<p>The network has the goal to provide data associated with liver cancer. These data include microarray data like aCGH, methylation, or gene expression. Further, imaging data like tissue microarrays and clinical data are considered.</p>
Weighting	high

Feature	Explanation
Number	G4
Name	Analyze data
Description	<p>Data of participating projects have to be made available for cross-project analyses.</p>
Weighting	high

Feature	Explanation
Number	G5
Name	Control data access and usage
Description	Data of SFB/TRR77 have to be protected. Depending on the type of data, only specific collaborators of the research network, all members of the network, or the general public can access data.
Weighting	high

Requirements of the research network

Feature	Explanation
Number	R1
Name	Create data
Description	Within the research network, microarray data on liver cancer specimen are produced. These data are not created within the IT platform itself, but distributed. Individual projects generate data for their own research. If required, these data are shared within the research network. This leads to the additional requirement of preparing the data for distribution.
Weighting	high

Feature	Explanation
Number	R2
Name	Retrieve external data
Description	Genomic data created within the network have to be annotated in terms of functionality and roles in regulatory pathways. This has to be done by correlating data bases available over the Internet like gene ontology or KEGG.
Weighting	medium

Feature	Explanation
Number	R3
Name	Represent data
Description	Data of the research network have to be easily accessible and locatable to enable their correlation. Thus, a central point of data access is required.
Weighting	high

Feature	Explanation
Number	R4
Name	Define syntax
Description	Syntax of the data available via the IT platform of the network has to be defined and documented in a suitable way. For data and analytical services, XML documents according to the WSDL specification are required.
Weighting	high

Feature	Explanation
Number	R5
Name	Define data model
Description	For all data accessible via the central IT platform data models have to be specified and documented. This is a prerequisite for data harmonization. For certain aspects documentation according to the UML-XMI specification is required.
Weighting	medium

Feature	Explanation
Number	R6
Name	Identify data
Description	All data sources need to be annotated with a unique URI to make them identifiable and accessible throughout the research network.
Weighting	low

Feature	Explanation
Number	R7
Name	Define semantics
Description	Semantics of the data provided for the SFB/TRR77 network have to be defined precisely by the data contributors. This is a prerequisite for cross-project data analysis.
Weighting	medium

Feature	Explanation
Number	R8
Name	Administrate intellectual property
Description	Projects of the research network require maintaining control over data they share via the IT platform. Further, suitable mechanisms for ensuring credit for data contributors have to be established.
Weighting	high

Feature	Explanation
Number	R9
Name	Protect data
Description	Data must be protected against unauthorized access. Since no identifying personal data are stored on the IT platform, this requirements helps mostly to enforce R8.
Weighting	medium

Feature	Explanation
Number	R10
Name	Show results
Description	Results of the network's data analyses have to be visualized in a central place. Different kinds of presentation, like tables or diagrams, have to be available.
Weighting	high

Feature	Explanation
Number	R11
Name	Integrate data
Description	Data of the research network have to be analyzed across project borders. Thus, they have to be integrated and made available for the whole network.
Weighting	medium

Feature	Explanation
Number	R12
Name	Define analytical methods
Description	For the correct analysis of the research network's data, specific biostatistical methods have to be selected and established. Flexibility is required to support a wide range of data types and alternative approaches.
Weighting	medium

Feature	Explanation
Number	R13
Name	Define analytical process
Description	The process of data analysis is comprised of one or more data sources and one or more analytical methods. These components have to be chained and connected to fulfill research tasks of the network.
Weighting	high

Feature	Explanation
Number	R14
Name	Static Workflow
Description	The IT platform has to provide predefined workflows for data analysis. Users have to be able to parameterize the workflow to run a specific analysis. Parameters could include gene symbols or statistical methods that can be chosen from a list.
Weighting	medium

Feature	Explanation
Number	R15
Name	Dynamic Workflow
Description	The IT platform has to provide tools for the users to design complete analytical workflows themselves. This includes selecting the available data and analytical services and chaining them into arbitrary pipelines.
Weighting	low