

UniversityHospital Heidelberg

# Coordination Centre for Clinical Trials (KKS)

**Structures and Functions** 

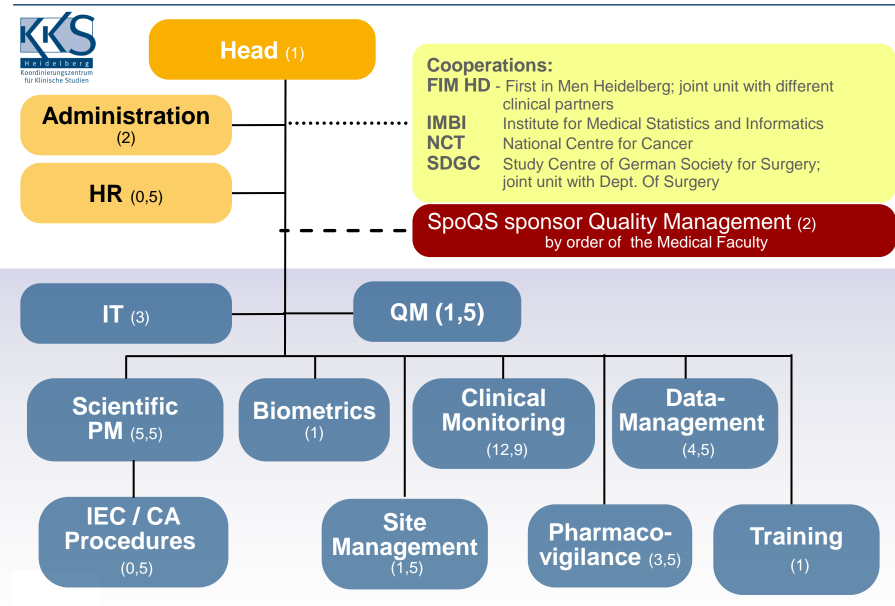


## Support provided by KKS

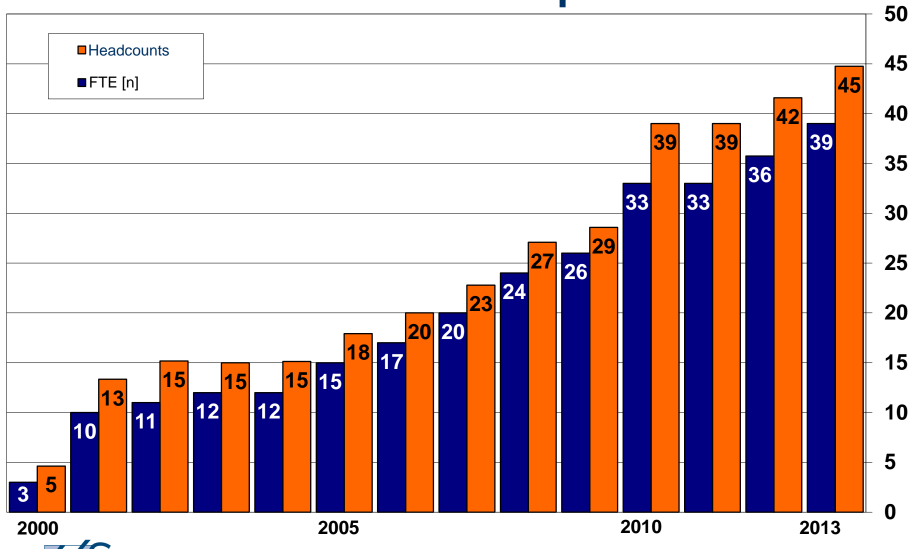
- Academic, non commercial trials
- Support of all aspects of clinical research
  - Sustainable, quality-assured study projects
- Optional services
  - For members of the University Hospital and for others
- Independence from University departments
  - No own academic ambitions
- Cost recovery



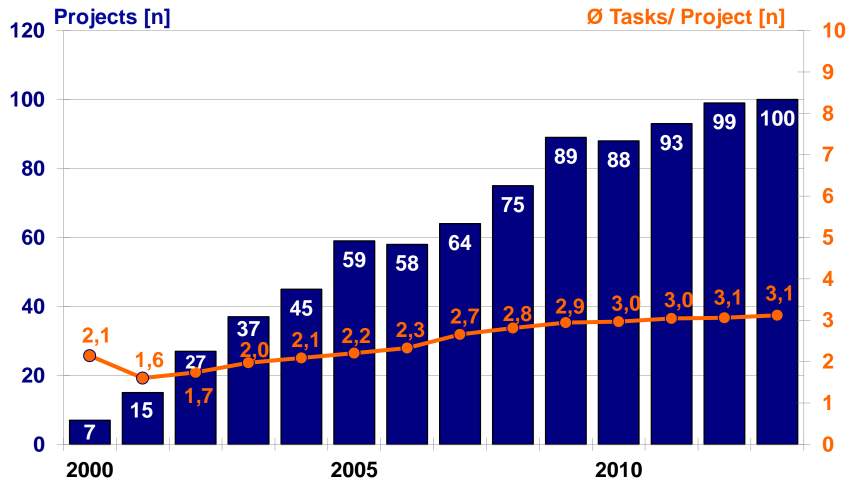
#### UniversityHospital Heidelberg



# **Personnel Development**



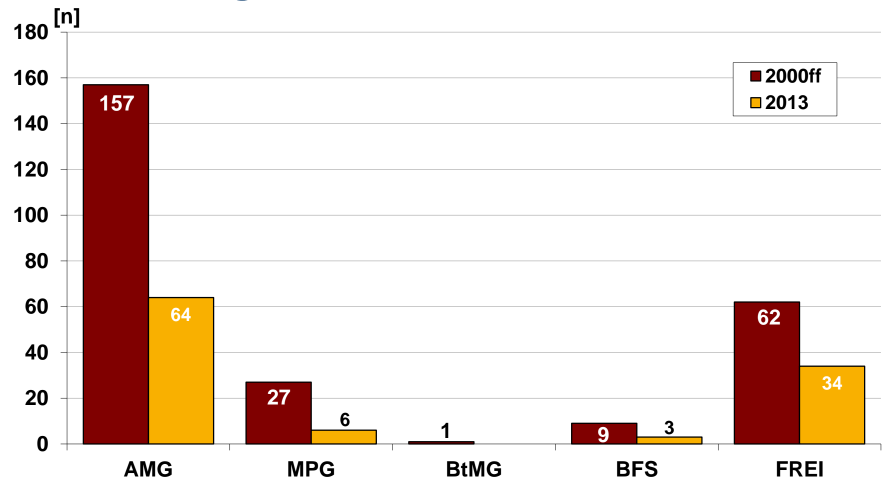
#### **Active Studies**





overall 238 studies since 2000

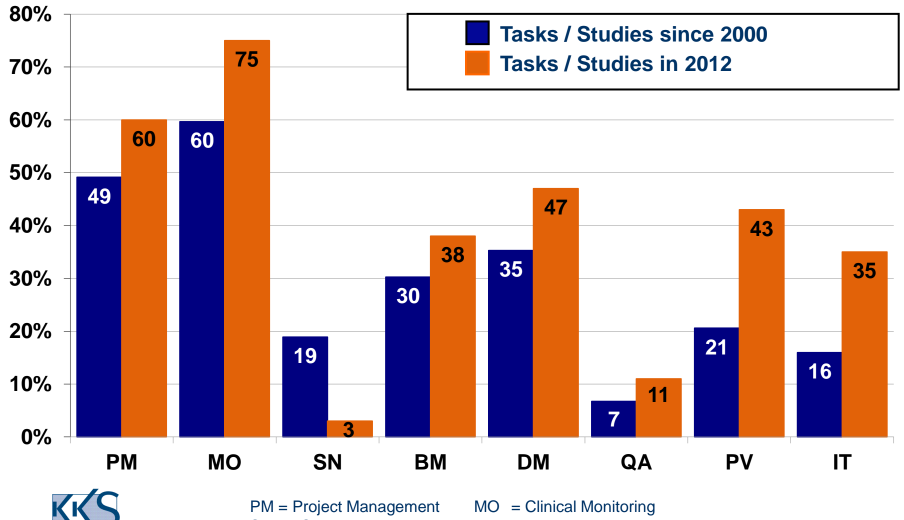
# **Legal Framework 2013\***





AMG = Arzneimittelgesetz (German Drug Law), MPG = Medizinproduktegesetz (German Medical Device Law), BtMG = Betäubungsmittelgesetz (Narcotics Law), BFS = Bundesamt für Strahlenschutz (German Federal Office for Radiation Protection), FREI = Not regulated by specific law

## Tasks Assumed by KKS

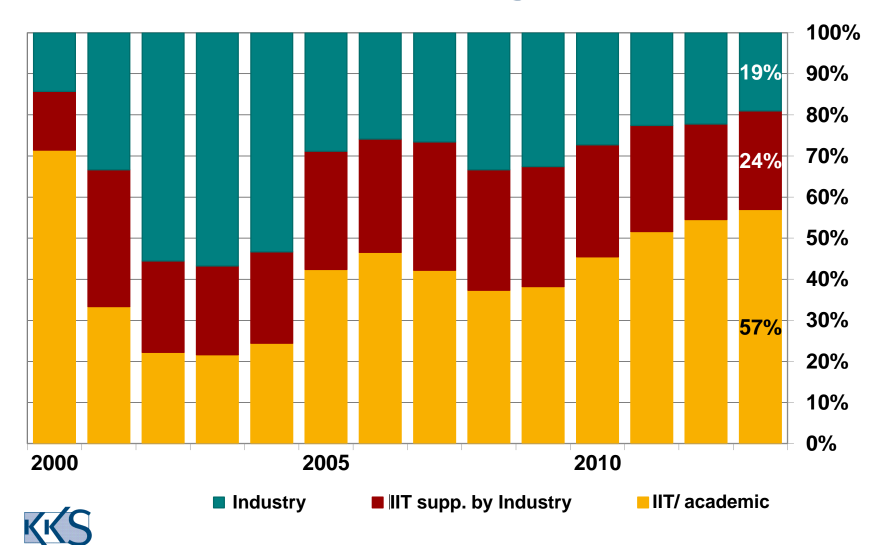


= Study Nurse Activities

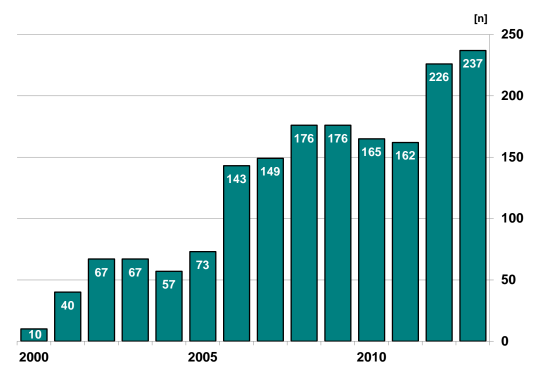
DM = Data Management = Pharmacovigilance BM = Biometry

QA = Quality Assurance

# **Funding**



#### **Consultations**



#### Expenditure 2013

- N=237
- ∑ 312 h, Ø 1.3 h,
   Several staff members (BM, PM, IT, QA etc.) involved, when necessary
- Specific services, such as providing templates, etc.

#### Compensation

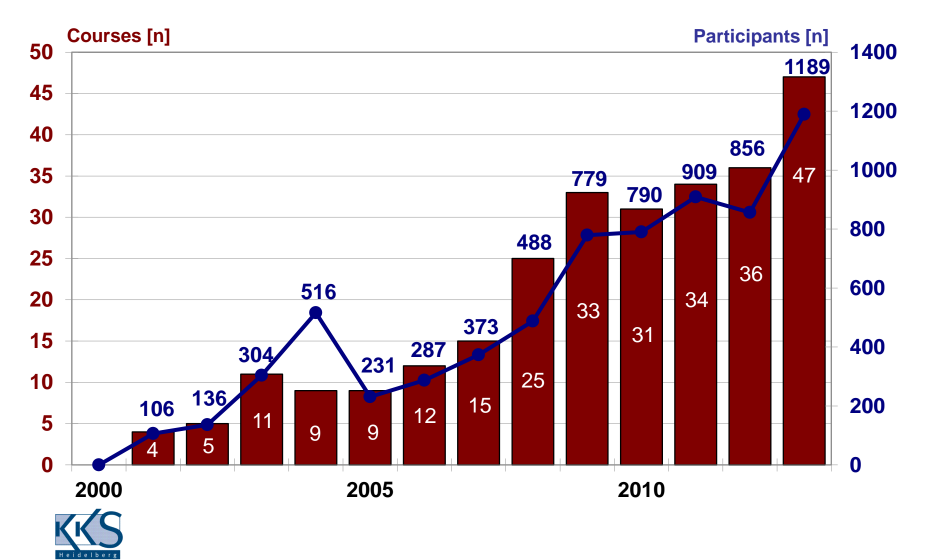
Generally free of charge

#### Contents

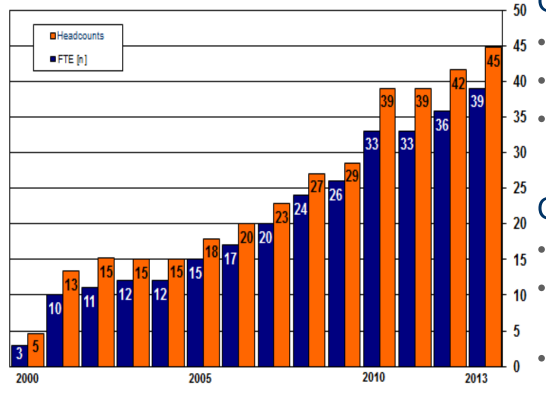
- General
  - AMG, insurance, etc.
- Project-related



# **Training Offers**



## **Personnel Development**



#### 50 Growth through

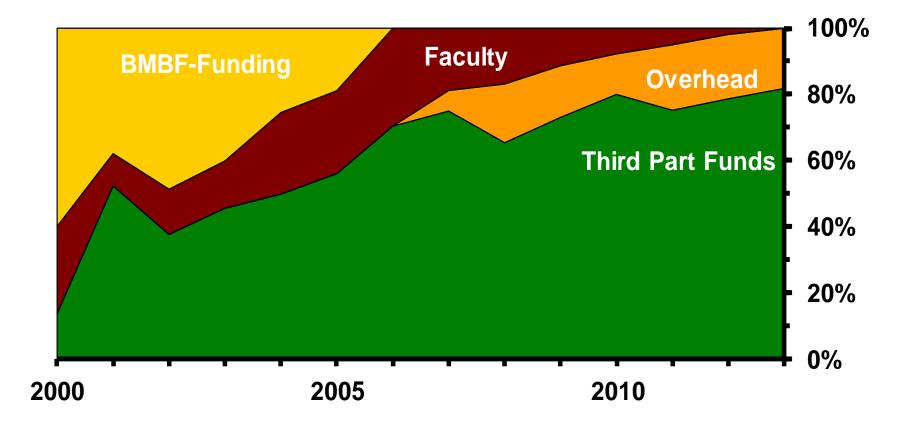
- Extension of existing tasks
- New tasks
- More comprehensive projects

#### Competence through

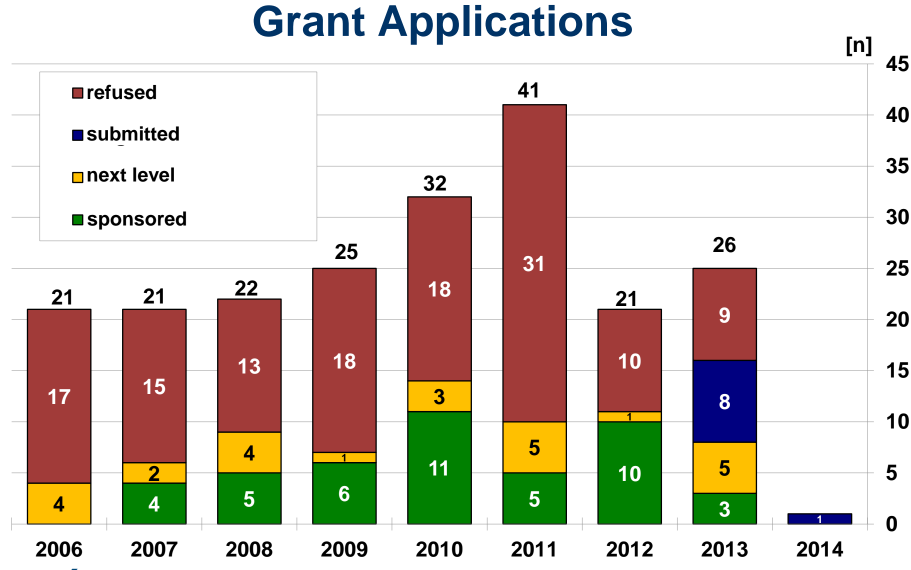
- Long-term employment
- Structured initial training and professional development
- Internal and external further education



# **Funding Shares**

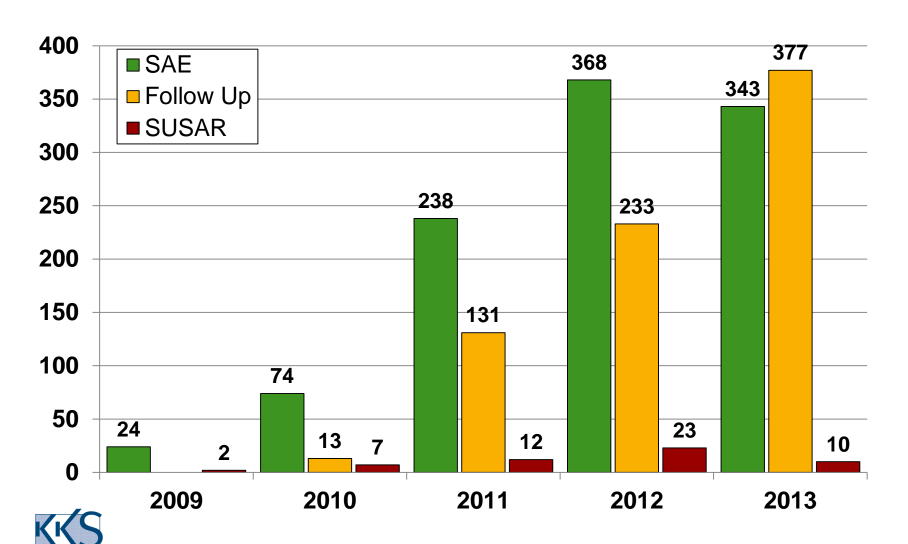








# **Safety Reports**



# **Clinical Monitoring / CRA**

- (on-site) support and monitoring of clinical trials
  - Adding to competence of investigators and coordinating investigators (LKPs)
  - Continuous quality control, adapted to each study and risk assessment

#### Objective

- Support of trial site
- Protection of trial subjects
- Best data quality and integrity
- Compliance with protocol and regulatory requirements

# Scientific Study Support / PM

- Unbureaucratic, confidential project consultation
- Study design and trial protocol development
- Project management
  - Ethics, regulatory affairs, (drug) safety
- Broad experience, independent of indication
  - AMG, MPG, non-AMG/ non-MPG, RöV, StrV, etc.
  - Comprehensive templates
- Document review
- Submission procedure to DFG, BMBF, EU etc.
- Consultation regarding research structure and contracts

# **Site Management Organization (SMO)**

- On-site support of the clinical trial
  - Scheduling of patient visits, conduct, documentation
  - Sample logistics
  - Preparation/ support of visits of (external) monitors and auditors
  - Updating of central documents (ISF)

#### Objective

- Well-planned and timely executed clinical trials
- Relief of medical personnel



## **Biometry**

- Conceptual consultation
  - Study design, sample size, etc.
- Biometric parts of the protocol
- Randomization procedures
- Statistical Analysis Plan, analyses
  - Sensitivity analyses for model assumptions and missing values
- Participation in the integrated final report



## **Data Management**

- Implementation of the clinical trial protocol
  - Variable lists
  - Case Report Forms (paper based, RDE)
- Data entry masks
  - Development, validation and data entry
  - Training for RDE systems
- Data Validation Plan (DVP)
  - Incl. query tracking
- Central randomization
- Support for the analysis



## **Quality Management**

- SOP system
  - Development, refinement and maintenance
  - Staff Training (incl. documentation)
  - Harmonized with the KKS Network and the TMF (Telematic Platform for Medical Research)
- Improvement of the staff's professional skills
  - On-the-job training and development plans
  - Internal and external training
- Consultancy services for trial centers
- QA review of essential documents
- Internal Audits



#### **IT Services**

- Validated systems for
  - Primary data entry in electronic format (RDE)
  - Data management
  - Statistical analyses
- Processes defined in SOPs
- Conceptual tasks regarding "IT-landscape" Clinical Research in Heidelberg

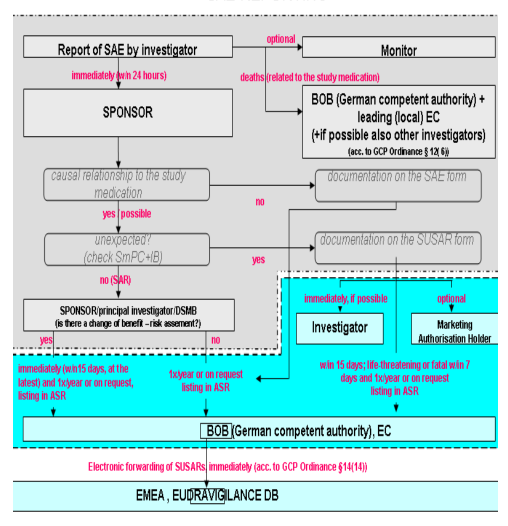


# **Pharmacovigilance**

#### **Main Tasks**

- Workflow Definition (Safety Manual)
- Preparation of study-specific data base
- SAE Management
- Follow-up of SAEs (obtaining of missing information)
- SUSAR Reporting to CA, IEC and study sites
- Preparation of Development Safety Update Reports (DSUR)

#### SAE-REPORTING





# **Training**

- Continual training offers
  - Impart practical knowledge
  - Qualification of attendees
  - Improvement of training situation in clinical research
- Curricula of several days or weeks for
  - Investigators
  - Study Nurses
  - Clinical Monitors
- Topical lectures and training
  - E.g. on recent amendment to the German Drug Law, drug safety, sponsor's role
- Lecture Series on Clinical Trials
  - Continual education as required by ICH GCP



## **Compulsory Consultation**

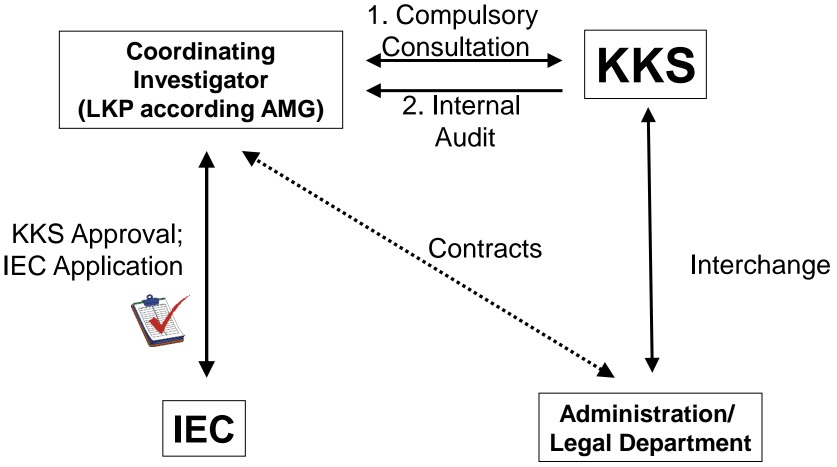
- Support the principal/coordinating investigator (LKP) in
  - 1. Designing clinical trials in compliance with German Drug Law, GCP Ordinance, data protection law, etc.
  - 2. Completely define the responsibilities associated with the clinical trial
  - 3. Avoiding funding shortfall
- Protection of the faculty's interests

#### Indirect

- Synergetic effects for parallel projects
- Enhancement of the reputation and competitiveness of clinical research



# Supervision of the Sponsor's Activities





# **Auditing**

- Support for external audits and inspections
- Recommendations for corrective actions
- Audits on behalf of Medical Faculty
  - Focus: trials regulated by the German Drug Law with principal/coordinating investigator in Heidelberg
  - Supervision of the sponsor's activities



# **FIM Heidelberg**

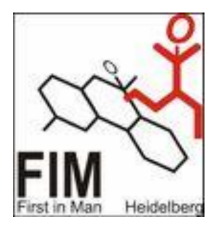
#### First in Man Studies ...

- "One stop shopping" strategy (planing, conducting and reporting)
- Multidisciplinary
  - KLIPS (Phase I Study Site on Clinical Pharmacology)
  - KKS (Coordination Centre for Clinical Trials)
  - IMBI (Medical Biometrics and Statistics)
  - NCT, DKFZ (pre- and clinical oncology)
  - Further clinical partners
- Contact

http://fim-hd.kks-hd.de

Email: fim@med.uni-heidelberg.de





## (Inter)national Activities

#### **National**

- KKS Network
  Coordinating Centres for Clinical Trials
- SDGC, Chir-Net
- TMF (German meta-organisation for networked medical research)
- Competence Networks

#### International

• European network





#### **Activities**



#### Interaction with funding organisations, currently regarding

- Pay scale grouping in grant applications
  - TV-Ä\*, TV-L<sup>#</sup>
- Applicability of GCP in psychotherapy trials
- Financing plans for clinical trials
  - Overhead costs, full costs
- Review process and statement for grant notification
  - E.g. Integrated Research and Treatment Centers (IFB):
     43 applications, only 3 accepted, reasons?

#### Exchange with ministries

- Comments on draft laws
  - Currently: German Radiation Protection Ordinance, X-Ray Ordinance, Drug Guidelines (AMR)



<sup>\*</sup>Sector pay scale for University Hospitals

<sup>#</sup> Public Sector Collective Agreement on Länder

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