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List of Publications

a) **Books etc.** (Editor, book chapters and teaching games)

- [1] J.-M. Arrang, M. Garbarg, J.-C. Schwartz, R. Lipp, H. Stark, W. Schunack, and J.-M. Lecomte.
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Autoreceptors and Heteroreceptors Evidenced by Histamine H₃ Receptor Ligands.
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- [7] R. D. E. Sewell, R. A. Glennon, M. Dukat, H. Stark, W. Schunack, and P. G. Strange.
Neurotransmitters, Agonists and Antagonists.
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(ed. H. J. Smith), Harwood Academic Publishers, Amsterdam 1998, pp. 387-433 (Text book).
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Neurotransmitters, Agonists and Antagonists.
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In: Histamine Research in the New Millennium (eds. T. Watanabe, H. Timmerman, and K.
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b) Articles (Original articles and reviews)

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GLISTEN/DFG Research Training Group, Erlangen/Germany (April 6-8, 2016).
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Editor-in-Chief

Archiv der Pharmazie – Chemistry in Life Sciences (2004 – heute)

Guest Editor of Hot-Topic Issue: *Penetration of Blood-Brain Barrier*
Current Medicinal Chemistry – Central Nervous System Agents 2002, Vol. 2(3), 75 pp.

Guest Editor with M. Schubert-Zsilavecz *Pharmazie in unserer Zeit* -
- Parkinsontherapeutika 2006, Vol. 35(3),
- Antitussiva und Expektorantien 2007, Vol. 36(6).

Guest Editor with T. Dingeramn *Pharmakon- Arzneimittel in Wissenschaft und Praxis*
- Aufmerksamkeitsdefizit-/Hyperaktivitätsstörung (ADHS) 2014, Vol. 2(1)

Editorial (Advisory) Board Member

Current Medicinal Chemistry (2006 - heute)

Current Medicinal Chemistry – Central Nervous System Agents (title change in 2006)
Central Nervous System Agents in Medicinal Chemistry (2000 – heute)

Die Pharmazie (2004 – heute)

Facta Universitatis, series Working and Living Environmental Protection (2014-heute)

Letters in Drug Design & Discovery (2012 – heute)

Medicinal Chemistry (2005 – heute)

Medicinal Chemistry Reviews – Online (2004-2005) Publikation eingestellt

Online Medicinal Chemistry (Open Access; 2007 - heute)

Open Medicinal Chemistry Journal (Open Access; 2007 - heute)

Recent Patent Reviews on CNS Drug Discovery (2006 - heute)

Organisation of congresses etc.

3rd European Graduate Student Meeting 2001, Member of local committee (February 23 – 25, 2001), DPhG, Frankfurt am Main

Frankfurter Forschungsreise – Kleine Moleküle mit großer Wirkung –
Das BIOZENTRUM im „Jahr der Lebenswissenschaften“ (5. Oktober 2001),
Organisation, Moderation und Führungen, Frankfurt am Main

4th European Graduate Student Meeting 2002, Member of local committee and local organizers (February 8 – 10, 2002), DPhG, Frankfurt am Main

Tag der Naturwissenschaften, Johann Wolfgang Goethe-Universität, BIOZENTRUM – Studiengang Pharmazie Frankfurt am Main.

(6. und 7. März 2002), Organisation, Führungen und Moderation,
dito (19. und 20. März 2003), Organisation, Führungen und Moderation.
dito (21. und 22. September 2005), Vortrag und Führungen.

Radiopharmaceutical Chemistry / Radiopharmacy, postgraduate diploma course (PDC), organised by the Universities of Frankfurt am Main, Leipzig and ETH Zürich, Course module: *Pharmacy I and Legislation*

- February 24 – March 7, 2003, co-organizer Frankfurt am Main
- September – October 2005, Course Director, Frankfurt am Main,
- September 2007, Course Director, Frankfurt am Main

90 Jahre Johann Wolfgang Goethe-Universität, *Pharmazie an der Goethe-Universität*, Festveranstaltung (29. Juni 2004), Frankfurt am Main, Organisation und Moderation

Frontiers in Medicinal Chemistry, Fachgruppentagung der DPhG (Pharmazeutische / Medizinische Chemie) und der GDCh (Medizinische Chemie), March 12 – 15, 2006, Frankfurt am Main/Germany, scientific committee (chair) and local organizing committee (chair)

XXXVIIIth Annual Meeting of the European Histamine Research Society and 20. Jahrestagung AVE Allergieverein in Europa e.V., May 13-16, 2009, Fulda/Germany, organizing committee.

BIOMEDCHEM on Histamine H₄ Receptors, International ESF-COST BM0806 Meeting WG4, Frankfurt/Germany, October 6, 2009, scientific committee (chair) and local organizing committee (chair).

GPCR – AN UPDATE ON STRUCTURE AND FUNCTION 2011. International Symposium for 21st Friedrich Merz-Guest-Professorship, Frankfurt/Germany, May 26, 2011, scientific committee (chair) and local organizer (chair).

PyMOL Schrodinger-Workshop Demo & Training, Frankfurt/Germany, May 9, 2012, co-organizer.

The future of H₄R research: the power of interaction, COST-MC and WG1-4 Meeting, Malaga/Spain, November 16-18, 2012, scientific committee.

Advances in Histamine H₄R Research, COST BM0806 Final Conference, MC & WG1-4 Meeting, Cape Sounion - Athens/Greece, March 21-23, 2013, scientific committee.

42nd Annual Meeting of the European Histamine Research Society (EHRS), Lodz/Poland, May 8-11, 2013, scientific committee.

Innovation in Cell Analysis – Zellanalytik-Seminar 2014, Düsseldorf/Germany, September 24, 2014, co-organizer, chair.

Workshop: *Chemical Drawing, Scientific Recherche and more*. Düsseldorf/Germany, November 12, 2014, organizer, chair.

Workshop: *Microscale Thermophoresis*. Düsseldorf/Germany, January 22, 2015, organizer, chair.

Annual Meeting of the German Pharmaceutical Society (DPhG), Düsseldorf/Germany September 23 – 25, 2015, organizer, chairman, scientific committee, organizing committee incl. Historical Presymposium 22.09.2015 and Postsymposium on Pain Management 26.09.2015.