Professor Neumann has three publications in journals of the Lancet group: one in Lancet 1991 and one in Lancet Oncology 2014.

The publication in 1991 was the first epidemiologic estimation of the prevalence of von Hippel-Lindau disease (approx. 1:40,000).

This publication showed in addition, that there are 2 types of von Hippel-Lindau disease, one type with pheochromocytoma and one type without pheochromocytoma. This has become international recognition and is still today the basis for classification of von Hippel-Lindau disease.

The publication of 2014 demonstrates that the concept of conservative adrenal surgery for pheochromocytoma is effective and the standard for pheochromocytoma in patients with multiple endocrine neoplasia type 2. This was a multicenter, multinational study involving 30 centers with more than 50 specialists. It demonstrates that Addisonian disease is a frequent longterm complication in patients with hereditary pheochromocytoma. This study is a key paper for organ sparing surgery in patients with hereditary tumor syndromes.

In the third publication we describe the complex challenge of adequate clinical and molecular genetic diagnosis of pheochromocytomas and paragangliomas as a review.


   Outcomes of adrenal sparing surgery or total adrenalectomy in pheochromocytoma associated with multiple endocrine neoplasia type 2: an international retrospective population based study
   The Lancet Oncology, epub ahead of print April 2014

2. Erlic Z, Neumann HPH.
   Diagnosing patients with hereditary paraganglial tumours.

3. Neumann HPH, Wiestler OD
   Clustering of features of Von Hippel-Lindau syndrome: evidence for a complex genetic locus
   The Lancet 1991;337:1052-1054

Hartmut Neumann