Dupuytren Society: joining forces in fighting Dupuytren's disease

Dupuytren Society was founded in 2003 in Germany. Goals are spreading knowledge of therapeutical options to treat Morbus Dupuytren and Ledderhose, support of patients and patient groups, supporting research, and – last not least – bringing medical doctors and patients together. Dupuytren Society considers itself a joint interest group of patients and doctors. Because Dupuytren's disease is not confined by borders Dupuytren Society has been acting internationally from the very beginning.

Today Dupuytren Society manages English and German web sites and runs two forums with close to 1,500 registered users from all over the world. Many more unregistered users are accessing our forum, typically 500 – 700 per day. We have been supporting several research projects, including this conference.