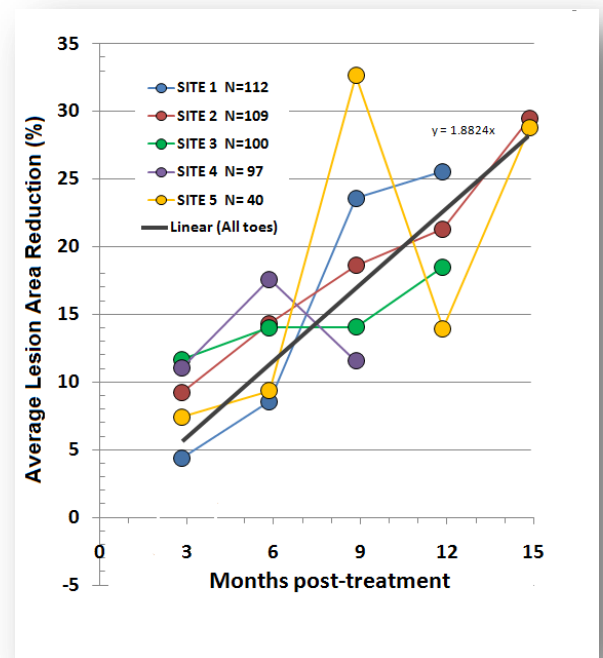


PINPOINTE FOOTLASER Preliminary Multi-Site Private Practice Retrospective Study Summary Demonstrates Steady Reduction in Lesion Size for Patients with Onychomycosis

This multi-site retrospective study was conducted in five private practices in the UK and the USA on 458 great toes from 265 sequential patients with onychomycosis that received a single 2-pass treatment with the PinPointe FootLaser. Before and after photographs collected by the doctors were computer analyzed using FDA validated planimetry to determine the average rate of improvement in this large sample. Safety was determined by analysis of patient records.

- There is continuous improvement in nail appearance to at least 15 months. The graph shows averages for each of five private practices.
- Statistical tests of efficacy are highly significant ($P < 0.001$).
- This large sample includes sequential patients and results include those patients with no or minimal treatment response.
- Sites report that many patients with complete resolution are not included because they were lost to follow-up.
- 71.4% of patients experienced continuous improvement in clear nail area.
- This study includes data submitted, in support of safety and efficacy, to US FDA, Health Canada, Therapeutic Goods Administration (Australia) and EU regulatory authorities to obtain needed clearances.
- The PinPointe FootLaser protocol is safe and effective.



Selected Cases - Before and After Photos

