

主な研究課題・発表代表論文

皮膚科学講座 Dermatology

研究領域 上皮情報解析医科学

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主な研究課題

- ・膿疱性乾癬の病態解明
- ・遺伝性皮膚・毛髪疾患の発症機構の解明
- ・形成外科領域の新規術式の開発に向けた研究

発表代表論文

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