

Letters

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Large-scale gene trapping in C57BL/6N mouse embryonic stem cells

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Errata

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^{OA}Open Access paper.



Cover False Color Photoshop image of *Salmonella enterica* serovar Typhimurium strain SL1344 grown on L agar for five days at 25°C, to show the rugose (lacey) phenotype. (Photo: Mark Roberts. [For details, see Thomson et al., pp. 1624–1637.])