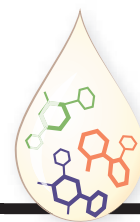


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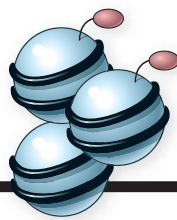
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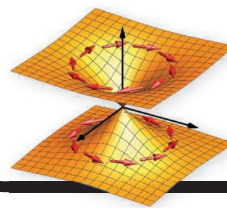


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Immunofluorescence microscopy identifies an intermediate cell in the cardiomyocyte lineage. An image of the heart of a 14.5-day mouse embryo shows this intermediate—progenitor derivatives expressing Hopx (red)—and the differentiated myocytes expressing troponin (green). *Jain et al.* demonstrate that Hopx-expressing cells promote cardiomyocyte commitment by coordinating signaling pathways in the progenitor niche. See page 1444 and dx.doi.org/10.1126/science.aaa6071. *Image: Epstein laboratory*

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