Reelin is a large extracellular glycoprotein involved in the development of architectonic patterns, particularly in the cerebral cortex and hippocampus, where it is synthesized primarily by Cajal–Retzius cells. In the hippocampus, Reelin also regulates the growth and/or distribution of afferent entorhinal and commissural axons. To assess further the possible action of Reelin on axonal growth, we used the three-dimensional collagen gel assay to measure axonal elongation from reeler cortical explants in the presence of Reelin. Because Reelin is proteolytically processed in vivo, normal explants and Reelin-transfected human embryonic kidney 293T cells were used, respectively, as sources of processed and full-length protein. The reliability of the assay was tested by demonstrating a clear repulsive action of semaphorin 3F (p < 0.0001). However, neither full-length nor processed Reelin exhibited any significant attraction or repulsion on cortical axons. Our results suggest that the reported effects of Reelin on axonal pathways are indirect, secondary to the architectonic disturbances that result from Reelin deficiency, and that the effects of Cajal–Retzius cells on connectivity are primarily independent of Reelin.

**Key words:** Reelin; axon guidance; collagen gel assay; cortical explants; hippocampus; Cajal–Retzius cells

Reelin, the extracellular glycoprotein defective in reeler mutant mice (D’Arcangelo et al., 1995), plays a key role in architectonic brain development (for review, see Lambert de Rouvroit and Goffinet, 1998). In the embryonic cortex and hippocampus, Reelin is synthesized by neurons in the marginal zone, including Cajal–Retzius (CR) cells, acts locally on end-migrating neurons of the cortical plate and instructs their radial organization. At the surface of target cells, Reelin binds to two lipoprotein receptors, very-low-density lipoprotein receptor (VLDLR) and apolipoprotein E receptor 2 (ApoER2), which relay the signal into the cell via the adapter Dab1 (Bar and Goffinet, 1998). In the embryonic cortex and hippocampus, Reelin-deficient mice are secondary to the brain malformation in reeler mice (D’Arcangelo et al., 1995), plays a key role in architectonic brain development (for review, see Lambert de Rouvroit and Goffinet, 1998). In the embryonic cortex and hippocampus, Reelin is synthesized by neurons in the marginal zone, including Cajal–Retzius (CR) cells, acts locally on end-migrating neurons of the cortical plate and instructs their radial organization. At the surface of target cells, Reelin binds to two lipoprotein receptors, very-low-density lipoprotein receptor (VLDLR) and apolipoprotein E receptor 2 (ApoER2), which relay the signal into the cell via the adapter Dab1 (Bar and Goffinet, 1998; Cooper and Howell, 1999; Trommsdorf et al., 1999). In addition to their Reelin-dependence on neuronal patterning, hippocampal CR cells were shown to guide entorhinal axons to the stratum lacunosum moleculare of the hippocampus, and perturbation experiments with the Reelin-blocking antibody CR50 suggested that Reelin may play a part in this guidance (Del Rio et al., 1997). However, observations that hippocampal afferents successfully reach their target in reeler mice, although with a significant delay (Borell et al., 1999a,b; Deller et al., 1999), suggest that other factors produced by CR cells are more important than Reelin, which could serve to promote collateral branching in terminal fields rather than guide entorhinal axons. Whether the developmental delay of hippocampal afferents in reeler mice reflects a direct action of Reelin on axonal growth or is secondary to the profuse architectonic malformation remains unclear. To study that question, the action of Reelin on cortical axonal growth was studied in vitro using the three-dimensional (3D) collagen gel assay. Because Reelin is cleaved in vivo, probably by a metalloproteinase (Lambert de Rouvroit et al., 1999), and the physiological consequences of this processing are unknown, the actions of both the full-length and the processed form of Reelin were analyzed. Neither form of the protein displayed a significant repulsive or attractive effect on axonal outgrowth. Although these observations cannot exclude a direct action of Reelin solely on a subset of axons, they strongly suggest that the observed disturbances of axonal growth in Reelin-deficient mice are secondary to the brain malformation in the mutant, possibly including disturbances of some recently described projections from CR cells (Ceranik et al., 1999).

**MATERIALS AND METHODS**

**Explant culture.** Normal BALB/c and homozygous reeler (Reln, Orleans allele) mice were used. Pregnancies were dated by checking females for the presence of a vaginal plug; the day of the plug was noted as embryonic day 0 (E0). Mice were killed by cervical dislocation and the embryonic brains were removed under cold anesthesia. Experiments were performed in accordance with National and Institutional Guidelines for animal care and were approved by the competent Animal Ethics Committee. To obtain cortical explants, the dorsal tiers of the hemispheres were dissected, meningeal membranes were peeled off, and the tissue was cut into 300 μm explants with a McIlwain tissue chopper (Campden Instruments, Leicester, UK).

Explants (E14–E18) were cultured in three-dimensional collagen gels as described previously (Lumsden and Davies, 1983; Toran-Allerand, 1990). Briefly, explants were embedded in 20 μl of collagen [9 parts rat-tail collagen, 0.9 parts 10× Eagle’s medium with t-glutamine (Life Sciences, Orleans, France), and 0.01 parts 0.3 M NaOH were mixed and added dropwise to the collagen solution, resulting in a final concentration of 0.025 M NaOH]. Explants were cultured in an atmosphere of 95% air and 5% CO2 and incubated at 37°C. After 1 day in culture and 2 days in the three-dimensional collagen gel, explants were fixed with 4% paraformaldehyde in PBS for 1 hour at room temperature, washed 3 times with PBS, and then permeabilized by 0.1% Triton X-100 in PBS for 10 minutes. Explants were then washed in PBS and incubated in blocking buffer (PBS with 3% normal goat serum and 0.5% Triton X-100) for 30 minutes. Explants were then incubated with primary antibodies in blocking buffer overnight at 4°C. The next day, after a 1-hour wash in blocking buffer, explants were incubated with secondary antibodies (goat anti-rabbit and donkey anti-goat Alexa Fluor 488 and Alexa Fluor 594, respectively) for 1 hour at room temperature.

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http://www.jneurosci.org/cgi/content/full/5827
Technologies, Grand Island, NY), and 0.1 parts 0.08 mM NaHCO₃ in 0.1
m NaOH) in culture dishes. The explants were positioned at a distance of 300–500 μm from each other or from cell pellets. The dishes were then
placed in an incubator at 37°C for 30 min to gelfy the collagen, before being covered with culture medium [Eagle’s medium with l-glutamine,
0.1% penicillin–streptomycin, Fisher’s cocktail (1 mg/ml BSA, 1 mg/ml
transferrin, 100 mM sodium selenite, and 250 μg/ml insulin), and 5% horse serum] as described previously (D’Arcangelo et al., 1997).

Reelin and semaphorin production in human embryonic kidney 293T
cells. Human embryonic kidney 293T cells were transfected with the
Reelin cDNA construct pC2 (D’Arcangelo et al., 1997), a Myc-tagged
semaphorin 3A construct, or a Myc-tagged semaphorin 3F construct
(provided by M. Tessier-Lavigne, University of California, San Fran-
cisco, CA). Cells were seeded at 3 × 10⁴ cells per 35 mm well and
transfected 16–20 hr later with 2 μg of Reelin cDNA using Lipofec-
tamine (10 μl in 1 ml of opti-MEM; Life Technologies). After 5 hr,
the supernatant and cell debris were removed and 2 ml of culture
medium was added (Iscoe’s modified Dulbecco’s medium with 10%
heat-inactivated fetal bovine serum, all from Life Technologies). Pellets
of transfected 293T cells were produced with the hanging-drop method
(Métin et al., 1997). At 12 hr after initiation of transfection, cell layers
were detached with trypsin, washed twice, and suspended in culture
medium with 1% serum (40 μl per 35 mm well). Drops (20 μl) of the cell
suspension were placed on the lids of 35 mm dishes, which were inverted
ded in medium containing 2 ml of medium. Hanging-drop cultures were
incubated for 14–16 hr, after which the cell pellets were harvested into
explant culture medium and embedded in collagen.

Quantification of axon growth. Axon growth was quantified by exami-
nation and photography under phase contrast or after immunostaining
with an anti-neurofilament 200 kDa antibody (clone RT97; Boehringer
Mannheim, Mannheim, Germany); analysis of results showed that both
methods were comparable. To avoid the possible effects of endogenous
Reelin, target explants were all from reeler mice. The zone surrounding
target explants was divided into quadrants, thus defining a proximal
and a distal quadrant in relation to the source of Reelin (either cell pellet or
normal explants), as schematized in Figure 1. The area covered by the
target explants was divided into quadrants, thus defining a proximal
and distal quadrants using Scion Image software (available from http://
rsb.info.nih.gov/nih-image). The ratios of the proximal and distal areas
were compared using Student’s t test after logarithmic transformation
and verification of homogeneity of variance (Bartlett test).

RESULTS
To verify that normal cortical explants and transfected 293T cells
expressed Reelin in the 3D collagen gel culture, supernatants were
tested by Western blot analysis with the G10 monoclonal
antibody, directed against the N-terminal region of Reelin (de
Bergeyck et al., 1997). As shown in Figure 2, transfected 293T
cells secreted a full-length Reelin of ~400–450 kDa, whereas
normal explants primarily produced processed Reelin, as evi-
denced by the demonstration of a predominant 180–200 kDa
N-terminal fragment. Control reeler explants (de Bergeyck et al.,
1998) and untransfected 293T cells did not secrete any Reelin.
These two situations were used to study the effect of full-length
and processed Reelin on axonal outgrowth. Similarly, the secre-
tion of semaphorin 3A and semaphorin 3F in the supernatants of
transfected cells was verified using anti-Myc antibodies. Both
semaphorin 3A and semaphorin 3F were detected as full-length
(95 kDa) and processed products, namely a 60 kDa C-terminal
fragment for semaphorin 3A and two fragments of 60 kDa and 35
dkDa for semaphorin 3F (Adams et al., 1997).

To assess the effect of Reelin on cortical axons, target reeler
explants were cultured next to normal cortical explants, used as a
source of processed Reelin, or next to a pellet of transfected 293T
cells, used as a source of full-length Reelin (Fig. 3). Recombinant
Reelin binds to both ApoER2 and VLDLR receptors and triggers
the phosphorylation of the Dab1 adapter, suggesting that it is
biologically active (Cooper and Howell, 1999; Trommsdorff et al.,
1999) (data not shown). In cultures, the control situation con-
sisted of two reeler explants or a reeler explant cultured next to
nontransfected cells. To get a positive control of the assay, the
effects of semaphorin 3A and semaphorin 3F were tested. After
3 d in vitro, axonal outgrowth from the reeler explant was anal-
ized as described above (Fig. 1). The following situations were
analyzed: 29 instances of reeler explants facing Reelin-
transfected 293T cells (Fig. 3A), 36 instances of reeler versus
normal explants (Fig. 3B), 15 control experiments of reeler–reeler
co-explants, 12 experiments with reeler explants facing nontrans-
fected 293T cells, 26 instances of normal explants facing sema-
phorin 3A-transfected cells (Fig. 3C), and 25 instances of normal
explants facing semaphorin 3F-transfected cells (Fig. 3D). The
results of the statistical analysis are shown in Table 1. Whereas a
significant repulsive action of semaphorin 3F on cortical axons
was evident (p < 0.0001), no repulsion was exerted on cortical
axons by semaphorin 3A in this system. The axonal outgrowth
from reeler explants was not statistically different when the expl-
ants faced sources of full-length Reelin (transfected cells),
sources of full-length Reelin (normal explants), or Reelin-
deficient sources (reeler explants or untransfected cells).
DISCUSSION
This study was undertaken following evidence that CR cells and possibly Reelin play a permissive role during the development of the entorhinohippocampal pathway (Del Rio et al., 1997). These findings suggested that, in addition to its well-known role in neuronal migration and architectonic patterning, Reelin may also regulate aspects of axonal growth and guidance (Gosh, 1997). The present results show that the outgrowth of axons from reeler cortical explants was not significantly influenced by the presence of full-length or proteolytically processed Reelin; therefore, Reelin does not appear to have any direct attractive or repulsive action on cortical growth cones, at least in vitro. The collagen gel assay (Lumsden and Davies, 1983) has been used successfully in various settings and its sensitivity is considered high (Métin et al., 1997; Chédotal et al., 1998). That the negative result obtained with Reelin is not attributable to a lack of sensitivity of the setup is also demonstrated by the clear-cut axonal repulsion exerted by semaphorin 3F. The somewhat unexpected absence of repulsion by semaphorin 3A may be related to the cells used for transfection, as indicated by the observation of different cleavage products for semaphorin 3A and semaphorin 3F (Adams et al., 1997).

The absence of detectable effects of Reelin shows that the protein does not exert any general direct influence on the guidance and growth of cortical axons. Thus far, Reelin has been shown to affect solely the distribution of commissural and entorhinal afferent axons to the hippocampus. Because the explant system examines outgrowth from several types of axons and thus is not selective, our data cannot rule out a direct effect of Reelin on a specific set of axons at a given developmental time. This reservation being made, our results are generally in agreement with other observations that suggest that the actions of Reelin on axonal pathways are likely indirect, secondary to its primary effect on architectonic patterning, and that the hodological effects of CR cells are primarily independent of Reelin. Studies in organotypic slice cultures showed that the ablation of CR cells prevents the ingrowth of entorhinal fibers. Interference with Reelin using the CR50 antibody has a less drastic effect: it does not prevent ingrowth of entorhinal fibers but reduces their branching and collateral expansion in the hippocampal target field (Del Rio et al., 1997; Borrell et al., 1999a). Similarly, entorhinal fibers reach their hippocampal target in Reelin-deficient mice, although with a significant delay, but they fail to form a normal contingent of collaterals and synapses (Borrell et al., 1999b). In addition, hippocampal commissural fibers in reeler mice disperse broadly in the stratum lacunosum moleculare, but they are strictly segregated in normal mice. This anomaly is correlated with the different distributions of CR cells and granule neurons in reeler versus normal mice (Borrell et al., 1999a,b; Deller et al., 1999). The observation that hippocampal CR cells project to the entorhinal cortex suggests that their processes may play a direct role in

Table 1. Results of coculture experiments

<table>
<thead>
<tr>
<th>Coculture condition</th>
<th>n</th>
<th>Mean log (P/D)</th>
<th>Result</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reeler/Reelin-transfected</td>
<td>29</td>
<td>-0.0147</td>
<td>NS</td>
</tr>
<tr>
<td>Reeler/normal</td>
<td>36</td>
<td>-0.1415</td>
<td>NS</td>
</tr>
<tr>
<td>Reeler/reeler</td>
<td>15</td>
<td>0.0967</td>
<td>NS</td>
</tr>
<tr>
<td>Reeler/untransfected cells</td>
<td>15</td>
<td>0.0297</td>
<td>NS</td>
</tr>
<tr>
<td>Normal/semaphorin 3A-transfected</td>
<td>26</td>
<td>-0.1566</td>
<td>NS</td>
</tr>
<tr>
<td>Normal/semaphorin 3F-transfected</td>
<td>25</td>
<td>-0.5267</td>
<td>p &lt; 0.0001</td>
</tr>
</tbody>
</table>

Data are analyzed by comparing axonal outgrowth in proximal (P) and distal (D) quadrants as shown in Figure 1 and described in Materials and Methods. n, Number of experiments; NS, not significant.

Figure 3. Examples of axonal growth in collagen gel culture assays. A, A Reeler explant (T) near a pellet of Reelin-transfected cells as the source (S). B, A Reeler explant (T) near a normal explant as the source (S). C, Normal explant (T) near a pellet of semaphorin 3A-transfected cells (S). D, Normal explant (T) near a pellet of semaphorin 3F-transfected cells (S). A repulsive action is clearly seen. Scale bar, 150 μm.
guiding entorhinal fibers to the hippocampal marginal zone. Surely, even if such actions are dependent on Reelin, they cannot be detected in collagen gel assays in which anatomical connections between explants are not present. Together, these studies suggest that the prominent effects of CR cells on the growth of entorhinal afferents to the hippocampus are not primarily attributable to Reelin; in addition, the present observation that Reelin has no direct effect on cortical growth cones provides a strong argument for this view. The consequences of Reelin deficiency observed in vivo are likely to be indirect, secondary to the profuse laminar malformation of the reeler entorhinal cortex and hippocampus. Given the possible role of the recently described projections from hippocampal CR cells to the entorhinal cortex (Ceranik et al., 1999), it would certainly be interesting to study that projection in reeler mice. Finally, the action of Reelin on collateral branching should be studied further, for example using in vitro systems such as slice cultures in which the architecture of the tissue is better preserved and can be more accurately assessed than in explant cultures.

REFERENCES