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Nemo regulates cell dynamics and represses the expression of *miple*, a midkine/pleiotrophin cytokine, during ommatidial rotation

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Abstract

Ommatidial rotation is one of the most important events for correct patterning of the *Drosophila* eye. Although several signaling pathways are involved in this process, few genes have been shown to specifically affect it. One of them is *nemo* (*nmo*), which encodes a MAP-like protein kinase that regulates the rate of rotation throughout the entire process, and serves as a link between core planar cell polarity (PCP) factors and the E-cadherin–catenin complex. To determine more precisely the role of *nmo* in ommatidial rotation, live-imaging analyses in *nmo* mutant and wild-type early pupal eye discs were performed. We demonstrate that ommatidial rotation is not a continuous process, and that rotating and non-rotating interommatidial cells are very dynamic. Our *in vivo* analyses also show that *nmo* regulates the speed of rotation and is required in cone cells for correct ommatidial rotation, and that these cells as well as interommatidial cells are less dynamic in *nmo* mutants. Furthermore, microarray analyses of *nmo* and wild-type larval eye discs led us to identify new genes and signaling pathways related to *nmo* function during this process. One of them, *miple*, encodes the *Drosophila* ortholog of the midkine/pleiotrophin secreted cytokines that are involved in cell migration processes. *miple* is highly up-regulated in *nmo* mutant discs. Indeed, phenotypic analyses reveal that *miple* overexpression leads to ommatidial rotation defects. Genetic interaction assays suggest that *miple* is signaling through Ptp99A, the *Drosophila* ortholog of the vertebrate midkine/pleiotrophin PTP receptor. Accordingly, we propose that one of the roles of Nmo during ommatidial rotation is to repress *miple* expression, which may in turn affect the dynamics in E-cadherin–catenin complexes.

Keywords

Nemo; Ommatidial rotation; Live-imaging; Gene expression; Miple; *Drosophila* eye

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Appendix A. Supporting information: Supplementary data associated with this article can be found in the online version at <http://dx.doi.org/10.1016/j.ydbio.2013.02.006>.

Introduction

The *Drosophila* adult eye is composed of around 800 units, or ommatidia, which are precisely oriented in mirror symmetric fashion relative to a dorsal–ventral midline, the equator. This pattern is generated during larval development in the eye imaginal disc, when ommatidial preclusters rotate 90° towards the equator adopting opposite chiral forms depending upon whether they lie dorsally or ventrally (Jenny, 2010). These patterning events closely follow a moving front of differentiation, the morphogenetic furrow (MF), which moves from posterior to anterior across the eye imaginal disc (Tomlinson and Ready, 1987). The Frizzled planar cell polarity (Fz-PCP) pathway controls the proper differentiation of R3 and R4 photoreceptors and, subsequently, the direction of ommatidial rotation (Seifert and Mlodzik, 2007). The direction of rotation depends on correct R3/R4 cell fate specification since misrotation is a common phenotype observed in loss- and gain-of-function mutants of PCP genes (Mlodzik, 1999). During this process ommatidial precursors rotate as a group, but independent of their undifferentiated, stationary neighbors, the interommatidial cells (IOCs) (Fiehler and Wolff, 2007). The exact cellular mechanisms that drive this behavior have not yet been established. In parallel to Fz-PCP signaling, which may regulate ommatidial rotation through effects on cytoskeletal elements via the Rho-Kinase Drok (Winter et al., 2001), this process is also regulated by the Epidermal growth factor receptor (Egfr) pathway (Brown and Freeman, 2003; Gaengel and Mlodzik, 2003; Strutt and Strutt, 2003). Egfr pathway members signal through both the Mitogen activated protein kinase (MAPK)/Pointed (Pnt) transcriptional cascade and Canoe (Cno) (Brown and Freeman 2003, Gaengel and Mlodzik, 2003), and also interact genetically with *E-cadherin* (*E-cad*) and *N-cadherin* (*N-cad*) during this process (Brown and Freeman, 2003; Gaengel and Mlodzik, 2003; Mirkovic and Mlodzik, 2006). Moreover, genes functionally related with cytoskeleton reorganization and cell adhesion act as downstream effectors of Egfr signaling, thus linking ommatidial rotation with cell adhesion and cytoskeleton rearrangements (Gaengel and Mlodzik, 2003; Mirkovic and Mlodzik, 2006). In addition, the cell adhesion molecules Echinoid (Ed) and Friend of Echinoid (Fred) are required at multiple steps during the ommatidial rotation process (Fetting et al., 2009), and Ed seems to be required to decrease Flamingo (one of the PCP core proteins) levels on non-rotating IOCs to permit correct rotation of ommatidial clusters (Ho et al., 2010). Other genes that have been shown to be required during ommatidial rotation are *nemo* (*nmo*), *scabrous* (*sca*) and *zipper* (*zip*) (Choi and Benzer, 1994; Chou and Chien, 2002; Escudero et al., 2007; Fiehler and Wolff, 2007, 2008; Mirkovic et al., 2011).

The *Drosophila* *nmo* gene encodes the founding member of the Nemo-like kinase (Nlk) subfamily of MAPKs (Brott et al., 1998). Nlk family members have regulatory roles in multiple developmental processes in vertebrates and invertebrates. Indeed, vertebrate NLK has been shown to participate in several signaling pathways, being activated by Transforming Growth Factor- β (TGF- β), Wnt, and IL-6 signaling (Brott et al., 1998; Ishitani et al., 1999; Kanei-Ishii et al., 2004; Kojima et al., 2005; Meneghini et al., 1999; Ohkawara et al., 2004), and to function downstream of nerve growth factor (NGF) (Ishitani et al., 2009). In addition, NLK phosphorylates and regulates the activity of several transcription factors in the nucleus such as T-cell factor (TCF)/Lymphoid enhancer factor (LEF), Signal transducer and activator of transcription 3 (STAT3), c-Myb, Smad4, the intracellular domain of Notch1 (Notch1-ICD) or Nuclear Factor- κ B (NF- κ B) through phosphorylation of its co-factor CREB binding protein (CBP) (Ishitani et al., 2010; Ishitani et al., 2003; Ishitani et al., 1999; Kanei-Ishii et al., 2004; Kojima et al., 2005; Meneghini et al., 1999; Ohkawara et al., 2004; Shi et al., 2010; Yasuda et al., 2004). In *Drosophila*, *nmo* is involved in diverse processes such as eye specification, synaptic growth, apoptosis, wing development, pair-rule patterning and circadian rhythms (Braid et al., 2010; Braid and Verheyen, 2008; Chiu et al., 2011; Merino et al., 2009; Mirkovic et al., 2002; Morillo et al., 2012; Verheyen et al., 2001;

Yu et al., 2011). Moreover, it seems that *nmo* mediates crosstalk between multiple signaling pathways since it antagonizes *Drosophila* Wg signaling (Zeng and Verheyen, 2004) and attenuates BMP signaling by phosphorylating Mad during wing development (Zeng et al., 2007). *Nmo* was originally identified as an ommatidial rotation-specific factor (Choi and Benzer, 1994), which was subsequently shown to be essential for regulating the rate of ommatidial rotation throughout the entire process (Fiehler and Wolff, 2008; Mirkovic et al., 2011). Genetic interaction assays suggested that *nmo* could be functionally related to the JNK cascade during ommatidial rotation (Fiehler and Wolff, 2008; Mihaly et al., 2001). Furthermore, it has been recently demonstrated that *nmo* genetically interacts with several core PCP components (*prickle*, *strabismus*), members of signaling pathways (*Notch*, *spitz*, *Egfr*) and genes encoding cell adhesion proteins such as E-cad (*shotgun*) and -catenin (*armadillo*) (Mirkovic et al., 2011). Indeed, it has been suggested that *Nmo* serves as a molecular link between core PCP factors and the E-cad- β -catenin (β -cat) complexes promoting cell motility during ommatidial rotation (Mirkovic et al., 2011).

In order to analyze more precisely the requirement of *Nmo* in the ommatidial rotation process, we used several strategies. In vivo analyses of wild-type and *nmo* mutant eye imaginal discs demonstrated that this gene regulates the speed of ommatidial rotation, as suggested from studies in fixed discs (Fiehler and Wolff, 2008). We also found that cone cell dynamics during this process is disturbed in *nmo* mutants and demonstrated that *Nmo* is required in these cells for correct ommatidial rotation. Our in vivo analyses also showed that interommatidial cells are less dynamic in *nmo* mutants than in wild-type discs. In addition, we performed a microarray study to identify genes that were deregulated in *nmo* mutant eye imaginal discs and that could be involved in ommatidial rotation. Four of the genes identified were validated and confirmed to be functionally linked to *nmo* by genetic interaction assays with several mutant alleles. In addition, phenotypic analyses revealed that the ommatidial rotation process is compromised when expression levels of some of those genes are modified. One of them is *miple*, which encodes a secreted heparin-binding protein that belongs to the midkine (MK)/pleiotrophin (PTN) family (Englund et al., 2006). In vertebrates, both MK and PTN are secreted cytokines that are implicated in many different processes, including cell migration (Muramatsu, 2010; Papadimitriou et al., 2009). Our results showed that *miple* overexpression causes rotation defects and that it interacts genetically with *nmo* and *nmo*-related genes, suggesting that *Nmo* is required to repress *miple* for correct ommatidial rotation.

Materials and methods

Fly stocks and genetics

Fly lines used in this study include: *nmo*^{PI} (Choi and Benzer, 1994), *sev* > *nmo* (Mirkovic et al., 2011), *aos*⁷ (Freeman et al., 1992), UAS-*miple* (Toledano-Katchalski et al., 2007), the *mthl8* allele *P{Mae-UAS.6.11}mthl8*^{F29.6} (Mukherjee et al., 2006), UAS-*Egfr*^{DN} (Freeman, 1996), *Egfr*^{CO} (Clifford and Schupbach, 1989). *ptp99A*¹, *shg*², *arm*⁴, *cut*-GAL4, *iRmiple*, *iRLRP1*, *iRAik*, UAS-*Dcr-2*, *P{EPgy2}CG32373*^{EY21017} (named in this paper as *EP*^{CG32373}) and the *unc-13-4A* overexpression line, *EP*^{EY04085} were obtained from the Bloomington stock center. *iRmthl8*, *iRCG32373* and *iRunc-13-4A* were obtained from the Vienna *Drosophila* RNAi Center. For UAS-*mthl8* transgenic lines full length *mthl8* cDNA LP02895 was subcloned into pUAST vector and flies were generated at Best-Gene Inc. (Chino Hills, USA.) by standard methods. Expression of several lines was checked by in situ hybridization with an *mthl8* probe in *en*-GAL4/UAS-*mthl8* embryos. *GMR* > *miple*, *armGFP*, *nmo*^{PI} and *cut*-GAL4, *nmo*^{PI} lines were generated by standard recombination methods. *nmo*^{DB}, FRT80 (Mirkovic et al., 2011) and *ey*-FLP; *ubiGFP*, FRT80 flies were used to induce mitotic recombination for *nmo*^{DB} clones analysis. *armGFP* was a gift of Silvia Muñoz-Descalzo (University of Cambridge, Cambridge, UK).

Live-imaging of pupal eye imaginal discs

Time-lapse imaging of pupal eye imaginal discs was performed as described (Escudero et al., 2007). Images were taken at 15 min intervals during at least 10 h in a Leica TCS SP confocal microscope. The images obtained were assembled and analyzed with ImageJ software. Measurements of IOCs apical areas were done manually with ImageJ. To quantify the number of IOCs disappearing in vivo during ommatidial rotation we followed each cell contained within the area comprised among 4 developing ommatidia from the beginning to the end of the process. IOCs that constricted their apical surface and subsequently disappeared were considered as dying cells. A total of 18 areas in *armGFP* control and 13 in *armGFP, nmo^{P1}* mutant discs were scored for this analysis.

Histology and immunohistochemistry

Analysis of adult retinæ was performed as previously described (Tomlinson and Ready, 1987). Sections were mounted in DPX and observed through the optical microscope in dark field. At least four eyes per genotype were analyzed. For ommatidial orientation analysis, the ImageJ angle measurement tool was used. Scanning electron microscopy analysis of adult eyes was performed following the critical point dry method (Wolff, 2011) using a Philips XL-30 microscope. For immunohistochemistry, eye imaginal discs were dissected and incubated for 20 min in 4% paraformaldehyde. Pictures were taken using a Leica TCS-NT confocal laser-scanning microscope. In the case of pupal retinæ, 42 h pupae were dissected and retinæ were stained as previously described (Bao and Cagan, 2005). Retinæ were mounted in Vectashield mounting medium (Vector) and pictures were taken in a Zeiss LSM510 microscope. The following primary antibodies were used: anti-dpERK (1:2000, Sigma, cat.# 8159) and anti-DECaD (1:10, DSHB DCAD2).

Microarray analysis

Total RNA was extracted from approximately 500 eye-antenna imaginal discs of synchronized L3 larvae and purified with the *mirVANA*TM miRNA isolation kit (Ambion#AM1260) following manufacturer's instructions. RNA quality analysis and quantification was performed in a Biorad Experion bioanalyzer. Three *armGFP* control and three *armGFP, nmo^{P1}* RNA samples were prepared and used to hybridize to Drosophila genome 2.0 Affymetrix microarrays following manufacturer's instructions (www.affymetrix.com) at the Multigenic Analysis Unit of the University of Valencia (Spain). Raw data reported in this paper have been submitted to Gene Expression Omnibus (Geo), accession GSE36127. Pre-processing of data was performed using the RMA (Robust Multi-Array) function of the *affy* package and differential expression analysis using the LIMMA (linear models for micro-array data) package, both from Bioconductor (www.bioconductor.com). For each gene the fold change was determined as the log₂ ratio of the two compared mean intensities, so that a fold change of 2 means a 2²-times increase in the expression of the corresponding gene in *nmo^{P1}* mutants. Adjustments for multiple testing were performed by using the Benjamini and Hochberg method (Benjamini, 1995). Only genes with adjusted *p-values* >0.05 were considered as positives.

RT-qPCR

One of the RNA samples used for the microarray analyses mentioned above and an independent one from *armGFP* and *armGFP, nmo^{P1}* eye imaginal discs were retrotranscribed to cDNA and used as template for RT-qPCR analyses. Total RNAs and cDNAs were also obtained from 3 independent samples of either *sev-GAL4* or *sev > Nmo* eye imaginal discs following the same procedure. Taqman technology was used for validation of all candidate genes by RT-qPCR but *nmo*, for which we used the sybr-green technology. Primers and probes (Table S1) were designed from a gene region as close as

possible to that corresponding to the microarray probes. Reactions using Taqman probes were performed in a LightCycler 480 II Real-Time PCR System (Roche Applied Science) following manufacturer's instructions. *-tubulin84B* was used as a reference gene for all the analyses. For *nmo* validation, a StepOne cycler (Applied Biosystems) was used.

Results

Live-imaging analyses of the ommatidial rotation process in wild-type and *nmo* mutant eye discs

Phenotypic analyses of flies homozygous for the *nmo^{PI}* hypomorphic allele showed that all ommatidia arrested at around 45° of rotation, which led to propose that rotation might occur in two 45° steps and that *nmo* would be required in the second step (Choi and Benzer, 1994). However, studies performed in stained imaginal discs homozygous for the same allele suggested that the ommatidial rotation rate in *nmo* mutants was lower than in wild-type discs throughout the entire rotation process (Fiehler and Wolff, 2008). Supporting this hypothesis, it was shown that most ommatidia in *nmo^{DB}* null mutant eyes failed to rotate at all and remained parallel to the equator (Mirkovic et al., 2011). Since *Nmo* overexpression caused ommatidial over-rotation, these data suggested that *Nmo* levels and/or activity directly correlated with the rate of rotation (Mirkovic et al., 2011). However, a deeper analysis of the cellular changes that occur during ommatidial rotation is necessary to understand how *nmo* is exerting its function in this process. In the last years, live-imaging techniques have been extensively used to determine how cells respond to patterning signals during development. In order to analyze in vivo potential differences in the cellular behavior during the ommatidial rotation process, we performed live-imaging analyses in wild-type and *nmo* mutant pupal eye imaginal discs (Escudero et al., 2007). An *armGFP, nmo^{PI}* recombinant line was generated to visualize in vivo ommatidial rotation in *nmo* mutants, and an *armGFP* line was used as a wild-type control. The experiments were performed with the *nmo^{PI}* hypomorphic allele since in our hands the null *nmo^{DB}* allele was lethal at early pupal stages. *armGFP* labels apical contours of epithelial cells and allows to visualize ommatidial preclusters and to determine their orientation and developmental stage (Fig. 1D–F). First, we confirmed that the *armGFP, nmo^{PI}* line reproduced the external adult phenotypes of *nmo^{PI}* mutants both in wings and eyes (data not shown). Wings were smaller than wild-type with a rounded morphology and showed a held-out phenotype (Choi and Benzer, 1994; Verheyen et al., 2001). Eyes were externally rough and narrower than wild-type (Choi and Benzer, 1994). Tangential sections of those eyes revealed a high number of under-rotated ommatidia (Fig. 1A, B), with a mean orientation angle of $59.3^\circ \pm 3.9^\circ$ with respect to the equator (Fig. 1C), thus reproducing the *nmo^{PI}* phenotype (Fiehler and Wolff, 2008).

White pupae of the corresponding genotypes were prepared and the cellular movements in eye imaginal discs were recorded (see Material and methods). Several observations could be made when analyzing the movies obtained from *armGFP* control discs. Our results showed for the first time in our knowledge that ommatidial rotation was not a continuous process, instead ommatidia moved forth and back until they reached their final orientation (rotation angle) (Movie 1). In addition, we demonstrated that cells in the ommatidial clusters rotate independently from the undifferentiated IOCs during this process, breaking and establishing new contacts with them (Movie 1). However, far from remaining static during the process, IOCs underwent clear shape changes independent of cell division, with continuous expansion and contraction of their apical surfaces (Fig. 1H and Movie 2). We also observed that some IOCs lying between developing ommatidial clusters, and usually not in contact with them, disappeared during the process suggesting they were suffering programmed cell death (Movie 3). We quantified the number of IOCs disappearing in an area delimited by four developing ommatidia finding that a mean of $1.5^\circ \pm 1.2$ IOCs disappeared per area (see Materials and Methods). Both apoptosis and apical cell shape changes have been

demonstrated to play important roles in the dynamics of developmental processes like embryonic dorsal closure by controlling forces that drive cell movements (Blanchard et al., 2010; David et al., 2010; Solon et al., 2009; Toyama et al., 2008). Similar analyses performed in *armGFP*, *nmo^{P1}* discs showed that although photoreceptor recruitment in such discs occurred as in *armGFP* controls, ommatidia rotated at a slower pace and they stopped rotating prematurely (Fig. 1G and Movie 4). This is the first in vivo demonstration of the role of *nmo* in regulating the ommatidial rotation rate during the entire process, as suggested from the studies in fixed eye imaginal discs (Fiehler and Wolff, 2008). We also observed that apical shape changes in the IOCs, and in turn changes in their areas, were less evident in *nmo^{P1}* discs than in controls during the process (Fig. 1H, compare Movies 2 and 5). Besides, we did not find IOCs disappearing in *nmo* mutant discs when performing similar analyses to those indicated above for control discs (0 ± 0 IOCs disappearing per area, *p*-value < 0.0001, compare Movies 3 and 5), which is consistent with suggestions that *nmo* plays a role in apoptosis in the embryonic epidermis and during pupal retinae development (Mirkovic et al., 2002). Taken together, our results suggest that apical shape changes and apoptosis of IOCs, together with the remodeling of their junctions with rotating cells, could contribute to the discontinuity of the rotation process. The reduction of apical shape changes in the IOCs observed in *nmo^{P1}* mutant disc when compared to controls could be a secondary effect of the reduced ommatidial movement in such discs. Alternatively, the reduction of IOCs dynamics and death could be contributing to disturb ommatidial rotation in *nmo* mutants.

The in vivo analysis of *nmo* mutant discs also revealed an abnormal behavior of the cone cell precursors, which rotate together with the photoreceptor precursors and independently of their undifferentiated neighbors, the IOCs (Fiehler and Wolff, 2007). We found that in control discs the equatorial and polar cone cell precursors break and establish new contacts with neighboring IOCs until they reach their correct position in the ommatidium (Fig. 2A and Movie 2). Quantitative analyses revealed that these cells typically break 2.7 ± 1.5 contacts and establish 2.2 ± 1.6 ($n = 12$) new contacts in *armGFP* discs over a period of ~ 7 h, encompassing from a stage in which R7 and anterior and posterior cone cell precursors have been recruited until equatorial and polar cone cell precursors reach their final position in the ommatidial cluster (Fig. 2C). However, these cells are significantly more static in *armGFP*, *nmo^{P1}* discs (Fig. 2B and Movie 5), in which they break an average of 0.3 ± 0.5 contacts and establish 0.5 ± 0.5 ($n = 10$) new contacts with neighboring IOCs (*p*-value < 0.005 in both cases) (Fig. 2C). These results suggest that the adhesive behavior of cone cell precursors is affected in *nmo* mutants, probably due to the reported activity of the Nmo kinase at the level of adherens junction complexes (Mirkovic et al., 2011). Next, we wondered whether this abnormal behavior of cone cells could have any consequences for the ommatidial rotation process. To address this question, we conducted a mosaic analysis by using the FRT/FLP system to generate clones of *nmo^{DB}* mutant cells in pupal eye discs. We measured the degree of rotation of mosaic ommatidia with a full complement of wild-type photoreceptors but with *nmo* mutant cone cells. This analysis showed that while completely wild-type ommatidia rotated over $88.9^\circ \pm 2.9^\circ$, lack of *nmo* function in at least one cone cell disrupted the ommatidial rotation process, with ommatidia remaining at $76.7^\circ \pm 6.1^\circ$ (Fig. 2D) (22 mosaic ommatidia in a total of 25 clones analyzed, *p*-value < 0.000001). These data indicate that *nmo* is also required in cone cells for correct rotation, and that the under-rotation phenotype in *nmo^{DB}* eyes is in part due to its lack of function in such cells. To confirm these results, we analyzed the ommatidial rotation angles in eyes from UAS-*nmo*/+; *cut-GAL4*, *nmo^{P1}/nmo^{P1}* flies, which expressed *nmo* specifically in cone cells with the *cut-GAL4* driver in a *nmo^{P1}* background, and compared them to *cut-GAL4*, *nmo^{P1}/nmo^{P1}* controls. The mean ommatidial rotation angle in *cut-GAL4*, *nmo^{P1}/nmo^{P1}* control eyes was significantly lower than in UAS-*nmo*/+; *cut-GAL4*, *nmo^{P1}/nmo^{P1}* eyes ($57.3^\circ \pm 3.4^\circ$ and $68^\circ \pm 3.2^\circ$, respectively; Fig. 2E). In addition we also observed that the percentage of ommatidia rotating over 50° was significantly higher in UAS-*nmo*/+; *cut-GAL4*, *nmo^{P1}/*

nmo^{P1} than in *cut-GAL4, nmo*^{P1}/*nmo*^{P1} controls (85.1 ± 2.2 % and 63.9 ± 3.6%, respectively; Fig. 2F). These results indicate that *nmo* expression in cone cells partially rescues the under-rotation phenotype of *nmo*^{P1} mutants. Taken together, our data demonstrate that *nmo* is required in cone cell precursors during ommatidial rotation.

Identification of genes differentially expressed in *nmo* mutant eye discs

The relationship of *nmo* to other genes and/or pathways that could explain its exact role during ommatidial rotation is still unknown. Genetic interaction assays with *bsk* and *TGF-activated kinase (dTak)* mutants indicated that *nmo* was functionally related to the JNK cascade in the eye (Fiehler and Wolff, 2008; Mihaly et al., 2001). In addition, genetic and biochemical studies have recently demonstrated a link between *nmo* and the Fz-PCP pathway, since Nmo physically interacts with the Stbm-Pk complex (Mirkovic et al., 2011). These experiments also showed that Nmo phosphorylates β -cat and E-cad, thus providing a potential mechanism by which Nmo could be influencing cell adhesion (Mirkovic et al., 2011). However, it has been also proposed that Nmo could regulate gene expression via its ability to phosphorylate several transcription factors and co-factors (Fiehler and Wolff, 2008). Thus, in order to identify new genes and/or pathways that could be related to *nmo* during ommatidial rotation we compared the expression profile of *nmo*^{P1} mutant eye imaginal discs to that of wild-type discs by using genome-wide microarray analyses. For doing so, total RNAs extracted from *armGFP* and *armGFP, nmo*^{P1} eye-antenna imaginal discs were used to generate cDNAs, hybridize Affymetrix *Drosophila* Genome 2.0 arrays (see Materials and methods) and analyze the expression profile of these genotypes. The analyses were performed with three independent RNA samples from each genotype. We thus identified 104 significantly up-regulated (50.7%) and 101 down-regulated genes (49.3%) (*adjusted p-value* < 0.05) in *nmo*^{P1} mutants with respect to controls (see Tables S2 and S3). As expected, *nmo* expression was significantly reduced in *nmo*^{P1} mutant discs (it was down-regulated 21.6-fold) (Fig. 3 and Table S3). Although the function of most of the genes identified is unknown, some participate in distinct biological processes related to the ommatidial rotation process, such as cell adhesion, signaling, cytoskeleton biogenesis/organization, and carbohydrate metabolism, involved in extracellular matrix biosynthesis (Tables S2 and S3). Since *nmo* seems to have a role in cell adhesion during ommatidial rotation (Mirkovic et al., 2011), we chose to focus on two up-regulated and two down-regulated genes for further analyses: *miple*, *methuselah-like 8 (mthl8)*, *unc-13-4A* and *CG32373*. Two main criteria were used to select these genes: (1) a high fold-change in their expression in *nmo*^{P1} mutant discs (*miple* and *mthl8* were up-regulated 19.6- and 19.3-fold, respectively; *unc-13-4A* and *CG32373* were down-regulated 5.8- and 8.3-fold, respectively) (Tables S2 and S3) and (2) their possible role in cell adhesion. *miple* encodes the *Drosophila* ortholog of the vertebrate MK/PTN cytokines (Englund et al., 2006). These secreted heparin-binding proteins are implicated in several processes, including enhancement of cell growth and survival, cell migration, angiogenesis and neurite growth (Muramatsu, 2010; Papadimitriou et al., 2009). In *Drosophila*, *miple* has a role in mesoderm spreading in the embryo during gastrulation (Toledano-Katchalski et al., 2007), a process that involves collective cell migration. In addition its expression has been shown to be regulated by the Egr ligand Spitz during eye development (Firth and Baker, 2007). *mthl8* encodes a G protein-coupled receptor that has been shown to interact in a two-hybrid assay with Thrombospondin, a protein that mediates adhesion through interaction with integrins (Chanana et al., 2007). Moreover, *mthl8* genetically interacts with members of the JAK/STAT pathway in the eye (Mukherjee et al., 2006). Little is known about the function of *unc-13-4A* and *CG32373*. *Unc-13-4A* has been shown to interact in a two-hybrid assay with Toutvelu (Stanyon et al., 2004), which participates in heparansulfate biosynthesis (Izumikawa et al., 2006; The et al., 1999) and was identified as one of the *Drosophila* orthologs of vertebrate proteins putatively implicated in neurotransmitter release (Lloyd et

al., 2000). Finally, *CG32373* encodes a protein containing an EGF-like calcium-binding conserved site and a Sushi/SCR/CCP domain, both involved in cell adhesion (de Vega et al., 2007; Nishimura et al., 2007). To confirm the microarray results for these genes, RT-qPCR analyses in *nmo* mutant and control eye discs were performed. In such experiments, we also analyzed *nmo* expression levels as a control. Our results showed that while *nmo*, *CG32373* and *unc-13-4A* are significantly down-regulated in the mutants, *miple* and *mthl8* are significantly up-regulated in the same individuals (Fig. 3), thus supporting the microarray results. The correspondence between the variations of *nmo* levels observed in the array and in the RT-qPCR analyses gave us a control of the reliability of the results. Therefore, we took the validated genes as candidates to be regulated by *nmo* during the ommatidial rotation process.

Candidate genes are functionally related to *nmo* and could have a role in the ommatidial rotation process

To determine the potential role of the candidate genes in ommatidial rotation or eye development we aimed to analyze the effect of their overexpression and RNAi with the *sev*-GAL4 and GMR-GAL4 drivers in an otherwise wild-type background. The results of these analyses are summarized in Table 1. First, we tested the *unc-13-4A* and *CG32373* genes, both down-regulated in *nmo^{P1}* mutant discs (Fig. 3), finding that while reduction of *CG32373* expression produced mild ommatidial rotation defects (Fig. 4E), as expected from the microarray results, no phenotype was observed when analyzing *unc-13-4A* RNAi (not shown). This result could indicate that *unc-13-4A* is not involved in ommatidial rotation. Alternatively, it could be that the *unc-13-4A* RNAi is not strong enough to reduce *unc-13-4A* expression to critical levels able to affect that process. According to expression results, neither *unc-13-4A* nor *CG32373* overexpression with EP lines had any consequence for eye development (not shown). Next, we analyzed the *mthl8* and *miple* genes, both up-regulated in *nmo^{P1}* mutant discs (Fig. 3). Consistently *miple*-overexpressing eyes displayed mainly ommatidial rotation defects and, less frequently, defects in photoreceptor specification (Fig. 4F). However, *mthl8* overexpression (see Materials and methods) had no effect on eye development (not shown), maybe because *mthl8* expression levels obtained with the transgenic lines are not high enough. When knocking-down *miple* and *mthl8* in the eye, we found that only in the last case ommatidial rotation defects were observed (Fig. 4D). Although this result may seem contradictory for the expression results, it has been reported that both over-expression and loss of function (LOF) of genes involved in rotation, such as components of Egfr signaling, give rise to ommatidial rotation defects (Gaengel and Mlodzik, 2003).

Next, to confirm the functional relationship between *nmo* and the validated genes, we performed genetic interaction assays by examining the effect of their overexpression and RNAi knockdown on the *sev > Nmo* eye phenotype (Fig. 4B), which is mainly due to ommatidial rotation defects and has been shown to be dosage sensitive (Fiehler and Wolff, 2008; Mirkovic et al., 2011). The results of these genetic interactions were obtained by quantifying the percentage of ommatidia with rotation defects in the corresponding genotypes (Fig. 4G) and are indicated in Table 1. We found that the four candidate genes interact genetically with *nmo*. In the case of *CG32373* and *miple*, the results of the assays were in agreement with the expression changes observed in *nmo^{P1}* mutants. The functional relationship of *miple* and *nmo* will be further confirmed and discussed below. However, the results obtained for the genetic interactions with *unc-13-4A* and *mthl8* were somehow contradictory (Fig. 4G and Table 1). To clarify this issue, we determined the expression levels of both genes in *sev > Nmo* eye discs by RT-qPCR, using *sev > GAL4/+* discs as controls (Fig. S1). Strikingly, we found that *mthl8* is highly upregulated in *sev > Nmo* discs, as it happened in *nmo* mutants. These results indicated that *mthl8* expression is dramatically

affected by either reduction or increase of *Nmo* function, which complicates the interpretation of the genetic interaction results. Indeed, we found that both LOF and *mthl8* overexpression enhanced the *sev* > *Nmo* phenotype (Fig. 4G). Since *mthl8* expression is almost undetectable in wild-type eye imaginal discs by in situ hybridization (Firth and Baker, 2007; Mukherjee et al., 2006), the results obtained in the RT-qPCR analyses confirmed that *mthl8* and *nmo* are functionally related, although the exact nature of this relation is difficult to establish. Regarding *unc-13-4A*, we found that the expression of this gene was not altered in *nmo*-overexpressing discs when compared to controls (Fig. S1), which does not give a clear explanation to the genetic interactions found between *unc-13-4A* and *nmo*.

Taken together, the results obtained in the genetic interaction assays with *sev* > *Nmo* as well as in the phenotypic analyses of flies overexpressing or with reduced expression of *miple*, *mthl8*, *unc-13-4A* and *CG32373* suggest a role of some these genes in ommatidial rotation. However, further analyses will be required to decipher their exact role(s) and to determine their functional relationship to *nmo* in this context.

Miple is functionally linked to *ptp99A* and with members of the E-cad– β -cat complex

As mentioned above, Miple is the *Drosophila* ortholog of the vertebrate MK/PTN cytokines, which participate in several processes involving cell migration (Muramatsu, 2010; Papadimitriou et al., 2009). These cytokines use different receptors to exert their function including PTP, Alk, LRP1, N-Syndecan and integrins 1–3 (Muramatsu, 2010; Papadimitriou et al., 2009). One of them, PTP, is a chondroitin sulfate proteoglycan that binds to both MK and PTN with high affinity and is recognized through the heparin-binding sites of these proteins. Its intracellular domain exhibits protein tyrosine (Tyr) phosphatase activity and has been shown to interact with β -cat in vertebrates to promote dephosphorylation of its Tyr residues (Meng et al., 2000). Indeed, it seems that PTN binding to this receptor inhibits the phosphatase activity of PTP, eventually inducing Tyr phosphorylation of β -cat and causing a disruption of the E-cad– β -cat complex (CCC) stability and cell adhesion (Meng et al., 2000; Perez-Pinera et al., 2006). Interestingly, *Nmo* can phosphorylate both Arm, the *Drosophila* ortholog of β -cat, and E-cad (Mirkovic et al., 2011). β -cat phosphorylation by *Nmo* occurs preferentially in three Ser/Thr residues in the C-terminal region of the protein. Although this phosphorylation did not affect CCC formation in vitro, it was shown to be biologically relevant for ommatidial rotation (Mirkovic et al., 2011). Considering this, and to determine whether the molecular mechanisms underlying Miple function during ommatidial rotation in *Drosophila* could be similar to those in vertebrates, we performed genetic interaction assays to identify candidate genes that could be acting downstream of *miple*. Although the *sev*-GAL4/UAS-*miple* flies displayed a clear rotation phenotype, it was externally too weak to be clearly modified. Therefore, we generated a *GMR* > *miple* recombinant line that showed a clear rough eye phenotype, which in tangential sections displayed rotation defects but mainly defects in photoreceptor recruitment (Fig. 5 A and not shown). The *GMR* > *miple* phenotype was dosage sensitive, as it was markedly suppressed by reducing *miple* expression with a UAS-IR*miple* line (Fig. 5A,B). Our results showed that dosage reduction of *nmo* with the *nmo*^{PI} and *nmo*^{DB} alleles was able to enhance the eye roughness of *GMR* > *miple* flies (Fig. 5C and data not shown), thus supporting the functional relationship between both genes. Next, mutant alleles of several candidate genes, including those encoding putative Miple receptors and downstream effectors as well as genes functionally related to *nmo* in the ommatidial rotation process were tested for interactions with *GMR* > *miple* (Fig. 5). Ptp99A was described as the putative *Drosophila* ortholog of the PTP receptor. Ptp99A is involved in motor axon guidance in the *Drosophila* embryo (Desai et al., 1996), although no phenotypic analyses have been performed in mutant larvae. Therefore, we wondered whether *ptp99A*

could be functionally linked to *miple* during eye development. Our results showed that reduction of *ptp99A* dosage (with the *ptp99A¹* allele) markedly suppressed the eye roughness of *GMR > miple* flies (Fig. 5D). Similar analyses with mutant alleles for genes encoding other putative Miple receptors such as *Alk* (Bazigou et al., 2007) and *CG33087*, which encodes the *Drosophila* ortholog of LRP1, yielded negative results (data not shown). To get further insight into the potential downstream effectors of *miple* function, we tested whether the *GMR > miple* eye phenotype was sensitive to endogenous levels of *arm* and *shotgun* (*shg*). Our results showed that the *arm⁴* and *shg²* null alleles were dominant suppressors of that phenotype (Fig. 5E,F), suggesting that Miple function is linked to components of the CCC. Signaling pathways downstream of vertebrate MK/PKN activation include MAPKs as important components (reviewed in Kadomatsu and Muramatsu, 2004). Consistent with this, *miple* overexpression activates MAPK during *Drosophila* embryonic mesoderm development (Toledano-Katchalski et al., 2007). Activation of Egfr signaling also leads to MAPK activation and this pathway is involved in ommatidial rotation (Brown and Freeman, 2003; Gaengel and Mlodzik, 2003; Strutt and Strutt, 2003). Indeed, it has been suggested that *nmo* could be regulating the rate of rotation through the Egfr pathway (Brown and Freeman, 2003; Choi and Benzer, 1994; Gaengel and Mlodzik, 2003; Mirkovic et al., 2011). To check for a potential relationship between *miple* and the Egfr pathway during rotation, we first analyzed MAPK activation by dp-ERK staining in *GMR > miple* eye imaginal discs but these experiments did not provide consistent results due to high signal variability both in discs overexpressing Miple and in controls (data not shown). We therefore tested for genetic interactions between *GMR > miple* and components of the Egfr pathway. We found that the *GMR > miple* eye phenotype was suppressed by down-regulation of Egfr signaling, both by expressing a dominant negative form of the receptor (*Egfr^{DN}*) (Fig. 5G) or with the *Egfr^{CO}* mutant allele (data not shown), and enhanced when up-regulating Egfr signaling by dosage reduction of the *aos* gene (using the *aos⁷* allele, Fig. 5I).

To determine whether all the observed genetic interactions were relevant for the ommatidial rotation process, we subsequently repeated the experiments with the *sev > miple* line but only testing the interacting alleles. In this case, we analyzed tangential sections of eyes with the corresponding genotypes by measuring ommatidial rotation angles (Fig. 5J). These analyses confirmed the genetic interactions found between *miple* and *nmo*, *ptp99A* and the CCC components, since dosage reduction of these genes was able to modify the ommatidial rotation phenotype of *sev > miple* eyes (Fig. 5J). However, we found that components of the Egfr pathway did not significantly modify the ommatidial rotation phenotype of *sev > miple* eyes (Fig. 5J), thus suggesting that the interactions observed when using the *GMR > miple* line could be affecting photoreceptor recruitment, a process that has also been shown to be regulated by Egfr and is altered in the *GMR > miple* line. It would be interesting in the future to investigate this possibility.

Taken together, the results obtained in the genetic interaction assays confirmed the functional relationship between *miple* and *nmo*. Moreover, the interactions found between *miple* and *arm* and *shg* are in agreement with the previous results in which null alleles of both genes were strong enhancers of the *sev > Nmo* phenotype (Mirkovic et al., 2011). As expected from the micro-array results the two genes interact genetically with *nmo* and *miple* in opposite directions, thus confirming that *Nmo* is required to inhibit *miple* expression. Our data also suggest that *Ptp99A* could be acting as a Miple receptor during eye development and points to a conservation of the MK/PTN signaling mechanisms between *Drosophila* and vertebrates. Interestingly, we also observed that the *ptp99A¹* allele significantly enhanced the ommatidial rotation phenotype of *sev > Nmo* (Fig. 4G), confirming that these genes are functionally linked during the process.

Discussion

Multicellular movements are essential in multiple morphogenetic processes. Among them, ommatidial rotation (OR) in the *Drosophila* eye is an example of a highly coordinated cell motility process, which is necessary to achieve the regular arrangement of retinal cells. The Nmo kinase is an important player during the entire rotation process, probably regulating the activity of the E-cadherin-catenin complex as well as integrating signals from several pathways such as Fz-PCP, N and Egr (Mirkovic et al., 2011). Here we demonstrate that Nmo is required in and regulates cone cell dynamics during OR, and that it could be also modulating IOC's death during the process. In addition, we have identified new OR genes whose expression is dependent on *nmo* activity, thus discovering new molecular mechanisms and regulatory pathways operating downstream of *nmo* during the process.

Live imaging reveals cone cell requirements of *nmo*

Our live-imaging analyses of pupal eye imaginal discs demonstrate that almost the complete OR process can be tracked, thus making it possible to analyze the behavior of individual cell types involved in OR. These analyses revealed for the first time that OR is not a continuous process. It was previously reported that ommatidial clusters, which contain photoreceptor and cone cell precursors, move independently of the undifferentiated stationary IOCs (Fiehler and Wolff, 2008). Thus intercellular contacts between both subsets of cells likely need to be constantly remodeled to enable OR without disrupting the integrity of the epithelium, as suggested (Fiehler and Wolff, 2008). Our results indicate that ommatidial clusters move forth and back during the process, probably as a consequence of the constant remodeling of cell contacts between the preclusters and the stationary IOCs. Although it could be a secondary effect of the ommatidial clusters rotation, one possibility could be that the contractile movements of the IOCs might generate forces able to pull and push the rotating clusters. This is an interesting hypothesis that would be worth to check in the future. In addition, programmed cell death might provide part of the forces affecting OR, similar to what happens during embryonic dorsal closure (Toyama et al., 2008). Our in vivo analyses of pupal eye discs homozygous for the *nmo^{P1}* allele confirm that *nmo* is required throughout the entire OR process (Mirkovic et al., 2011). Consistent with this, we found that the rate of OR in *nmo^{P1}* mutants is lower than in controls at any point of the process. Our results suggest that *nmo* could be affecting OR through regulation of several distinct cellular aspects. First, we find that *nmo* regulates cone cell dynamics, which are very static when *nmo* function is reduced. In particular, the number of contacts they establish/break with surrounding cells in *nmo* mutant discs is significantly lower than in controls. Our mosaic analyses in pupal discs confirm that *nmo* is required in cone cells for correct OR. Moreover, the rotation defects in *nmo* mutants are partially rescued by expressing Nmo specifically in cone cells. Although a recent study already demonstrated a role of the cone cells in ommatidial rotation (Fetting et al., 2009), this is the first evidence of Nmo requirement in these cells during the process. Second, the absence or reduction of IOCs programmed cell death during OR in *nmo^{P1}* mutants suggests that Nmo is required to eliminate surplus cells and supports a dynamic role for apoptosis during this process. Finally, we also find a reduction of IOCs apical shape changes in *nmo* mutants with respect to controls. One possibility could be that this is a secondary effect of the reduced rate of OR in the mutants. However, an alternative hypothesis could be that *nmo* might regulate the contractility of these cells, and in turn the forces they are contributing during retinal development. Supporting this hypothesis, it has been reported that *nmo* regulates the activity of the CCC by directly phosphorylating β -cat (Mirkovic et al., 2011), and that *zip¹* suppresses the OR phenotype of *sev > Nmo* eyes (Fiehler and Wolff, 2008). These results suggest that Nmo could be acting upstream of the actin-myosin contractility by modulating polarized remodeling of adherens junctions (Mirkovic et al., 2011), and could support a role of Nmo

in regulating the adhesive properties of CCs, as suggested by its involvement in the dynamics of these cells.

A differential expression screen for Nmo targets

Our differential expression analyses in eye imaginal discs revealed that Nmo regulates the expression of several genes that encode cell adhesion and signaling molecules, among others. Preliminary data of four candidate genes (*miple*, *mthl8*, *unc13-4-A* and *CG32373*) indicate that some of them interact genetically with *nmo* and that their deregulation causes OR defects, supporting the validity of the microarray results. Interestingly, one gene identified in these analyses, *four wheel drive* (*fwd*), which was upregulated in *nmo^{PI}* mutant discs (see Table S2), has been recently isolated as a dominant modifier of a gain-of-function eye phenotype of the Fz-PCP core components Diego (Dgo) and Prickle (Pk) (Weber et al., 2012), further supporting a functional relationship between *nmo* and the Fz-PCP pathway (Mirkovic et al., 2011). However, in this study we have mainly focused on the analysis of *miple* function during OR. We demonstrate that *miple* overexpression leads to OR defects, consistent with the finding that it is highly up-regulated in *nmo^{PI}* mutant eye discs. Interestingly, one of the vertebrate orthologs of *miple*, PTN, is involved in the modification of cell adhesiveness (Perez-Pinera et al., 2006). Both PTN and MK, the second vertebrate ortholog of *miple*, contain a thrombospondin type I repeat homologous domain, and belong to the thrombospondin superfamily of adhesion molecules (Kilpelainen et al., 2000). In addition, the Mthl8 receptor has been shown to interact with Thrombospondin in a two hybrid assay (Giot et al., 2003), and two additional members of this family, *m-spondin* (*mspo*) and *fat-spondin*, were also identified in the microarray analyses as being significantly up- and down-regulated, respectively, in *nmo^{PI}* mutants (see Tables S2 and S3). Taken together, these results suggest that members of the thrombospondin superfamily could be important during OR and support the role of Nmo in regulating cell adhesion.

A possible role of miple in CCC regulation

To get further insight into the potential role of *miple* during OR, we tested whether its function was sensitive to endogenous levels of several candidate genes. We found that *miple* interacts genetically with *ptp99A*, which encodes the *Drosophila* ortholog of the PTP receptor, thus suggesting that *miple* signaling in *Drosophila* could be similar molecular mechanisms as its vertebrate counterparts. The genetic interaction between *miple* and the CCC members *arm* and *shg* also indicates that it could be participating in the remodeling of adherens junctions in the eye, as has been suggested in vertebrates (Perez-Pinera et al., 2006). The mechanism by which *miple* could affect OR remains however unclear. As described above, Nmo phosphorylates β -cat in Ser/Thr residues upon binding to PCP core components (Mirkovic et al., 2011), a process that could hinder phosphorylation of Tyr residues thus stabilizing the CCC (Fig. 6A). We propose that in absence of *nmo* function these Tyr residues would be exposed to phosphorylation leading to CCC destabilization. Interestingly, it has been shown that the PTP receptor in vertebrates is able to promote dephosphorylation of β -cat Tyr residues, and this activity is inhibited after PTN binding (Meng et al., 2000; Perez-Pinera et al., 2006). A similar situation could exist in the *Drosophila* eye. Here, in wild-type, where *miple* expression is repressed, β -cat would be phosphorylated by Nmo on Ser/Thr residues and this, in cooperation with Ptp99A activity, would lead to low levels of Arm phosphorylation in Tyr residues and to CCC stabilization (Fig. 6A). Upon *miple* up-regulation in *nmo* mutants, the phosphatase activity of Ptp99A could be inhibited, thus contributing to an increase of Tyr phosphorylation of Arm, to CCC destabilization (Fig. 6B) and in turn to rotation defects. Since it is unclear which Nmo expressing cells are also targets of Miple function and considering that ommatidial clusters rotate independently from the surrounding IOCs, this model could be probably applied to the interface between rotating and non-rotating cells. Alternatively, it has been proposed that

Nmo could regulate the rate of rotation independently of the PCP complexes through Egfr and/or N signaling (Mirkovic et al., 2011). Our results show however that *miple* does not interact with components of the Egfr pathway at the OR level thus discarding this pathway as a link between *miple* and *nmo* during this process.

Finally, an interesting question is how *miple* expression could be regulated by *nmo* in the *Drosophila* eye. Regarding this, it has been shown that MK expression in vertebrates is regulated by NF- κ B (You et al., 2008), a transcription factor whose activity is in turn negatively regulated by NLK, the vertebrate ortholog of Nmo, through phosphorylation of its co-factor CREB binding protein (CBP). Our preliminary results demonstrate that dosage reduction of *nejire* (*nej*), which encodes the *Drosophila* ortholog of CBP, dominantly modifies the eye phenotypes produced by overexpression of either *nmo* or *miple* (VM-S and NP, unpublished results). These data suggest a potential mechanism by which *nmo* could be regulating *miple* expression and would explain *miple* up-regulation in *nmo*^{P1} mutant eye discs.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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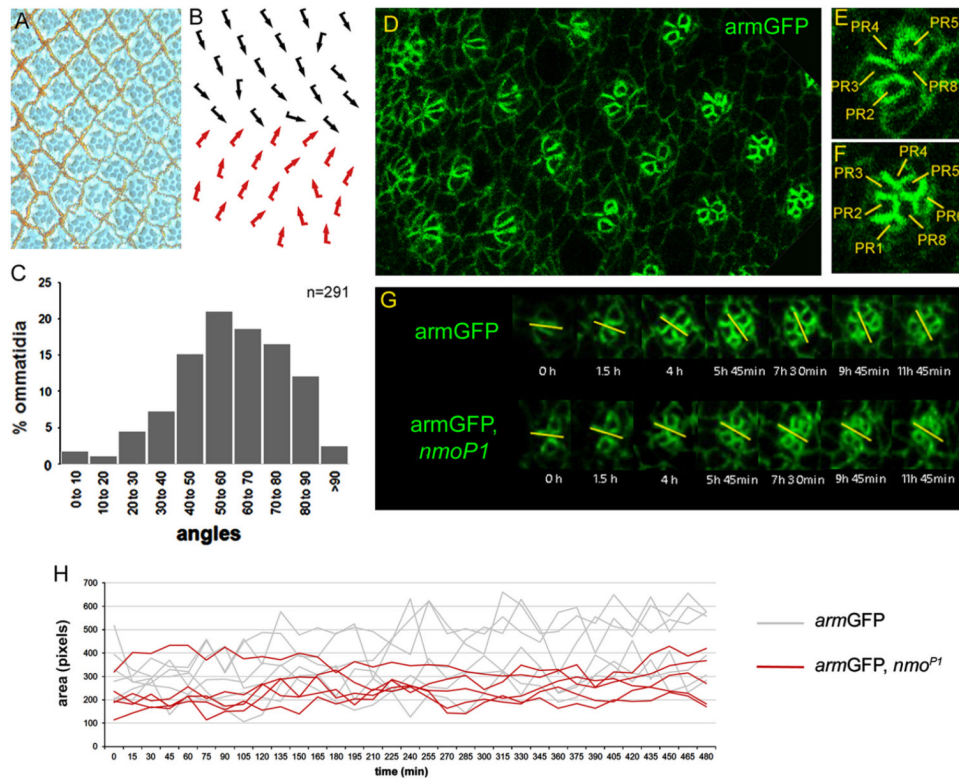
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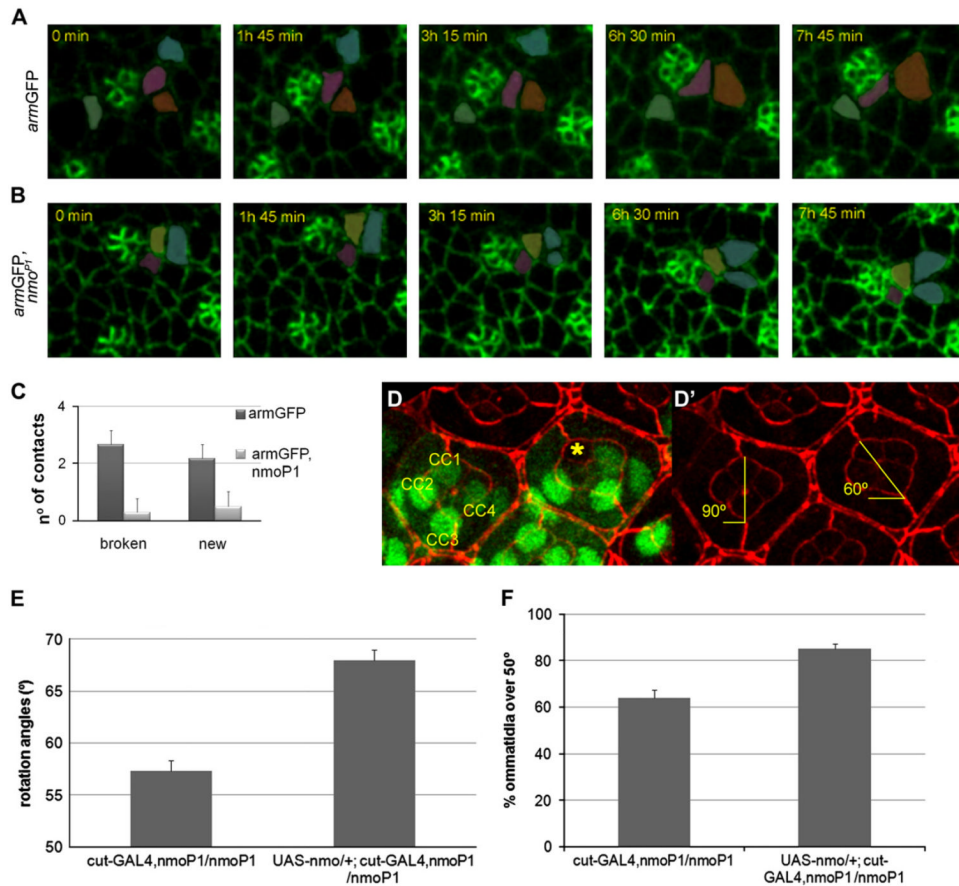
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**Fig. 1.**

Live-imaging analyses in eye imaginal discs reveal that *nmo* regulates the rate of ommatidial rotation and IOC's dynamics. (A–B) Tangential section of *armGFP*, *nmo^{P1}* adult eye (A) and the corresponding schematic representation with ommatidia arranged around the equator (B), with dorsal and ventral chiral forms indicated by black and red arrows, respectively. (C) Bar chart illustrating the percentage of ommatidia (*y*-axis) that are oriented at the angles indicated (*x*-axis) in *armGFP*, *nmo^{P1}* eyes, in which the most represented angles range from 50° to 70°. (D–F) *armGFP* protein localization in eye imaginal discs. A transgene with the adherens junction protein linked to GFP labels apical cell contours and outlines cell boundaries in an area of the eye imaginal disc posterior to the morphogenetic furrow (D). Magnified views of an ommatidial precluster that has initiated rotation (E), in which the five photoreceptor (PR) cells are labeled with their numbers, and an older one (F), in which almost all the PRs have been recruited. (G) Time-lapse series showing individual ommatidia during rotation after ~ 12 h from *armGFP* (upper panel) and *armGFP*, *nmo^{P1}* (lower panel) eye imaginal discs. The yellow bars mark the orientation angle of ommatidia with respect to the equator and the time on each photograph is referred to the first image of the series. The rotation rate of *armGFP*, *nmo^{P1}* ommatidia is slower than that of *armGFP* controls. (H) Quantification of several IOCs areas (number of pixels/cell) over time in *armGFP* (gray lines) and *armGFP*, *nmo^{P1}* (red lines) eye imaginal discs. Note that fluctuations of IOCs areas are sharper in wild-type controls than in *nmo* mutant discs, which is consistent with the observation that apical shape changes in IOCs are reduced in such mutants.

**Fig. 2.**

Nmo regulates cone cell dynamics during ommatidial rotation. (A–B) Time-lapse series showing the dynamics of equatorial cone cell precursors (marked in pink) in *armGFP* (A) and *armGFP, nmo^{P1}* (B) eye discs over the course of ~8 h. Time on each photograph is referred to the first image of the series. Other neighboring cells have been artificially colored to better follow the cell contacts. While in (A) the cone cell precursor breaks and establishes new contacts with neighboring cells, in (B) the cone cell precursor remains static on its initial position without breaking or forming new contacts. (C) Bar chart representing the number of cell contacts broken and established by the equatorial and polar cone cell precursors. (D–D') Confocal image of a 42 h pupal retina showing an ommatidium with wild-type cone cells (CC1 to CC4) and a mosaic ommatidium in which one of the cone cells (yellow asterisk) is mutant for *nmo^{DB}* (marked by the absence of GFP staining). In both ommatidia, the whole PR complement is wild type (not shown). In (D) GFP staining (green) marks *nmo⁺* cells, DE-Cad staining (red) shows orientation of ommatidia with respect to equator. In (D') only the DE-Cad staining is shown. The orientation angles of both ommatidia are marked in yellow. Loss of *nmo* function in one of the cone cells avoids complete ommatidial rotation (to 90°). (E–F) Overexpression of *nmo* specifically in the cone cells with the *cut-GAL4* driver partially rescues the *nmo^{P1}* rotation phenotype. Quantification of orientation angles in ommatidia from *UAS-nmo/+; cut-GAL4, nmo^{P1}/nmo^{P1}* and *cut-GAL4, nmo^{P1}/nmo^{P1}* eyes (E) reveals a significant increase of the mean angle orientation when *nmo* is overexpressed in cone cells. The percentage of ommatidia with an orientation angle over 50° also significantly increases in *UAS-nmo/+; cut-GAL4, nmo^{P1}/nmo^{P1}* eyes when compared to *cut-GAL4, nmo^{P1}/nmo^{P1}* controls (F).

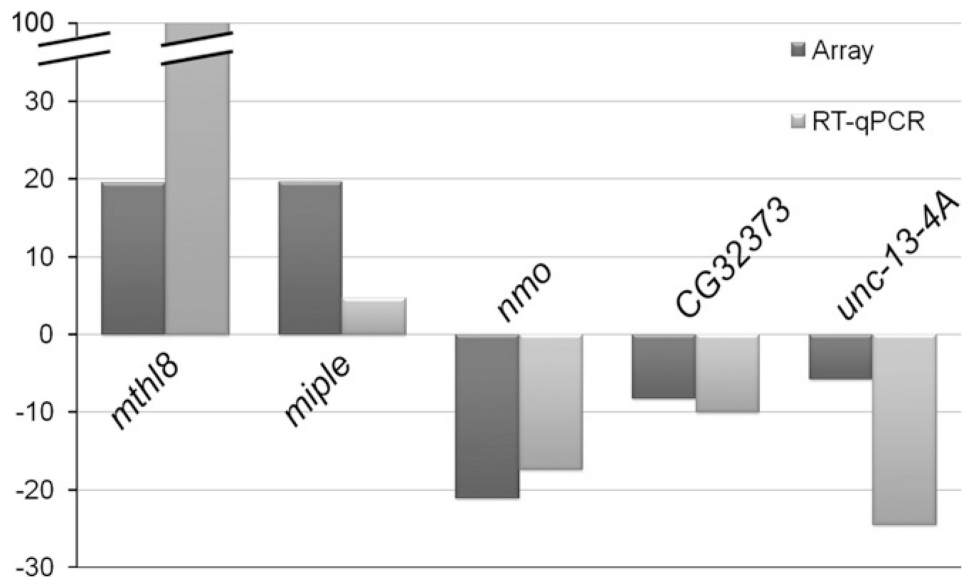


Fig. 3. Correlation of the differential expression of candidate genes in arrays and RT-qPCR. Graphic representation of fold changes in the expression levels of *nmo* and four candidate genes identified in the microarray analyses of *armGFP*, *nmo^{PI}* eye imaginal discs. Comparisons between the fold change values obtained in RT-qPCR and microarray analyses are shown.

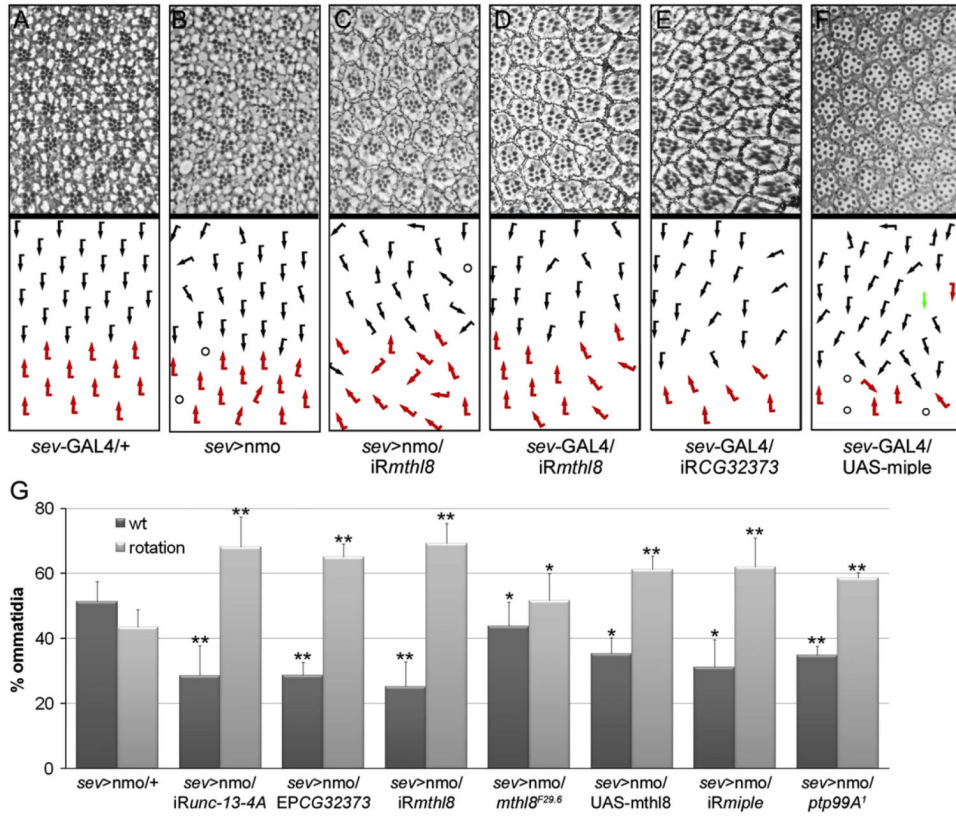


Fig. 4. Candidate genes interact genetically with *nmo* and could have a role in ommatidial rotation. (A–F) Tangential sections of adult eyes of the indicated genotypes and the corresponding schematic representations of ommatidial orientation with respect to equator, with dorsal and ventral chiral forms indicated by black and red arrows, respectively. Circles represent ommatidia with incorrect number of PRs and the green arrow indicates a symmetric ommatidium. Experiments were performed at 29 °C in (A), (D) and (E) and at 25 °C in (B), (C) and (F). (G) Quantification of the percentage of wild type ommatidia and ommatidia with rotation defects in adult eyes of the indicated genotypes. Note that in all cases, there is a significant modification of the *sev>nmo* phenotype (**p-value*<0.05, ***p-value*<0.01, Student's *t*-test).

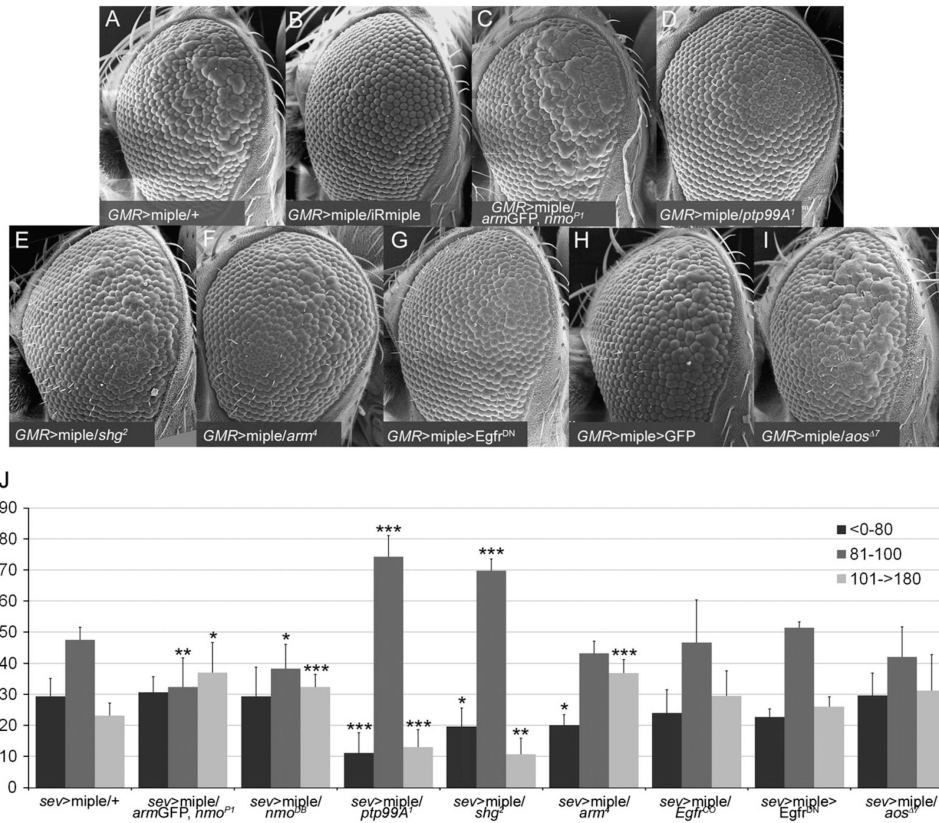


Fig. 5. *miple* interacts genetically with *nmo*, *ptp99A* and members of the CCC in the OR context. (A–I) Scanning electron microscope images of female adult eyes showing the external phenotype of *miple* overexpression with the *GMR*-GAL4 driver (*GMR > miple*). This phenotype (A) is dominantly suppressed by an UAS-IR *miple* line (B), the *ptp99A¹* (D), *shg²* (E) and *arm⁴* (F), and by down-regulation of Egfr signaling with a UAS-Egfr^{DN} transgene (G). In contrast, the *GMR > miple* phenotype is enhanced by *nmo^{P1}* (C) and *aos¹⁷* (I). Note that this phenotype is not modified by GFP overexpression (H). All experiments were performed at 25 °C. (J) Graphic representation of genetic interactions with the *sev > miple* ommatidial rotation phenotype. A quantification of the ommatidial orientation angles for each genotype is represented. The different angles have been grouped in three categories: > 0°–80° (under-rotated ommatidia), 81°–100° (wild type ommatidia) and 101°–> 180° (over-rotated ommatidia). Asterisks indicate statistically significant modification of the *sev > miple* phenotype for a given ommatidial orientation category (**p*-value<0.05, ***p*-value<0.01, ****p*-value<0.005, Student's *t*-test).

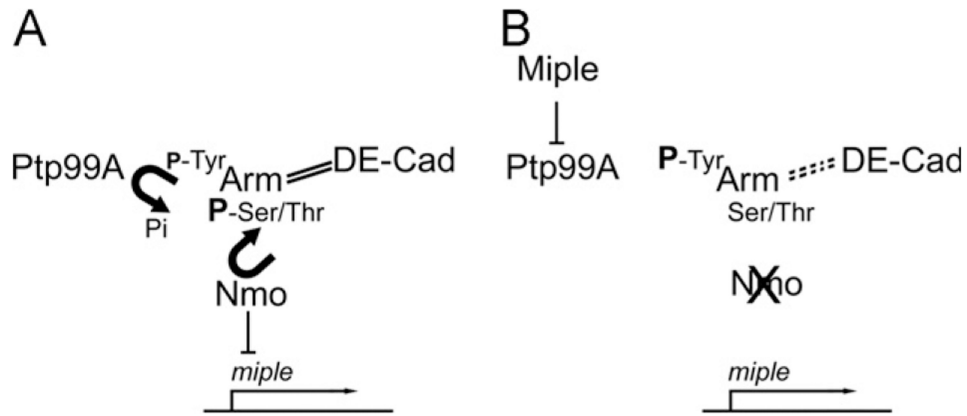


Fig. 6. Model for the effect of loss of *nmo* function on CCC destabilization mediated by Miple during ommatidial rotation. (A) In a wild-type situation, Nmo inhibits *miple* expression and phosphorylates Arm in Ser/Thr residues thus stabilizing the CCC and hindering phosphorylation of Tyr residues by other kinases. The levels of Tyr phosphorylation could be also lowered by the phosphatase activity of Ptp99A. (B) In *nmo* mutants, *miple* expression is activated and the Miple protein binds to the Ptp99A receptor, inhibiting its phosphatase activity against the Tyr residues of Arm and leading to CCC destabilization. Solid and dashed lines between Arm and DE-cad represent stabilization and destabilization of the complex, respectively. P represents phosphorylation of the corresponding amino acid, and its size correlates with phosphorylation levels.

Table 1Genetic interactions and phenotypic analyses of *nmo* targets.

Gene	<i>sev</i> -GAL4 GMR-GAL4		<i>sev</i> > <i>nmo</i>	
	OE	iRNA	OE	iRNA
<i>unc-13-4 A</i>	-	-		e
<i>CG32373</i>	-	+	e	
<i>mthl8</i>	-	+	e	e
<i>miple</i>	+	-		e

OE, overexpression; iRNA, RNA interference; -/+, wild type/ommatidial rotation phenotype; e/ , enhancement/no modification of the *sev* > *nmo* eye phenotype.