An ENU-Mutagenesis Screen in the Mouse: Identification of Novel Developmental Gene Functions

Carolien Wansleeben*a, Léon van Gurp, Harma Feitsma*b, Carla Kroon, Ester Rieter*c, Marlies Verberne, Victor Guryev, Edwin Cuppen, Frits Meijlink*

Hubrecht Institute, Royal Netherlands Academy of Arts and Sciences and University Medical Centre Utrecht, Utrecht, The Netherlands

Abstract

Background: Mutagenesis screens in the mouse have been proven useful for the identification of novel gene functions and generation of interesting mutant alleles. Here we describe a phenotype-based screen for recessive mutations affecting embryonic development.

Methodology/Principal Findings: Mice were mutagenized with N-ethyl-N-nitrosurea (ENU) and following incrossing the offspring, embryos were analyzed at embryonic day 10.5. Mutant phenotypes that arose in our screen include cardiac and nuchal edema, neural tube defects, situs inversus of the heart, posterior truncation and the absence of limbs and lungs. We isolated amongst others novel mutant alleles for Dll1, Ptprb, Plexin-B2, Fgf10, Wnt3a, Ncx1, ScribbScribb, Scribbled homolog [Drosophila]) and Sec24b. We found both nonsense alleles leading to severe protein truncations and mutants with single-amino acid substitutions that are informative at a molecular level. Novel findings include an ectopic neural tube in our Dll1 mutant and lung defects in the planar cell polarity mutants for Sec24b and Scribb.

Conclusions/Significance: Using a forward genetics approach, we have generated a number of novel mutant alleles that are linked to disturbed morphogenesis during development.

Introduction

In the developing embryo common molecular pathways are used to generate different cell types, tissues and organs. Although many of these pathways are reasonably well understood, not all of their components are known. Genetic approaches have been of decisive importance in discovery of these factors. Originally, genetic analysis depended on mutations occurring accidentally, but during the last two decades researchers have become less dependent on chance as reverse genetic approaches using targeted genetic analysis depended on mutations occurring accidentally, and reproduction in any medium, provided the original author and source are credited.

Copyright: © 2011 Wansleeben et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This work was supported by a grant from the Dutch government (BSIK program 03038, Stem Cells in Development and Disease) to FM. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: f.meijlink@hubrecht.eu

# Current address: Department of Cell Biology, Duke University Medical Center, Durham, North Carolina, United States of America

# Current address: Philips Research Labs, Eindhoven, The Netherlands

# Current address: Department of Cell Biology and Institute of Biomembranes, University Medical Centre Utrecht, Utrecht, The Netherlands

Citation: Wansleeben C, van Gurp L, Feitsma H, Kroon C, Rieter E, et al. (2011) An ENU-Mutagenesis Screen in the Mouse: Identification of Novel Developmental Gene Functions. PLoS ONE 6(4): e19357. doi:10.1371/journal.pone.0019357

Editor: Laszlo Orban, Temasek Life Sciences Laboratory, Singapore

Received December 31, 2010; Accepted March 31, 2011; Published April 29, 2011

PLoS ONE | www.plosone.org 1 April 2011 | Volume 6 | Issue 4 | e19357
mouse, it being a viviparous organism, is the choice one has to make of the embryonic stage of analysis. The seminal screen of Kasarskis et al. [10] focused on embryonic day 9.5 embryos, while in the screen by Herron et al. [11] letuses of E18.5 day were analyzed. While early-lethal mutations do not show up in the latter screen, in the former no mutants with phenotypes that become evident at later stages are detected. A different approach to further focusing on specific potential phenotypes was applied by Zarbalis et al. [5], who used disturbed expression of a Dcx5/6-LacZ transgene as screening criterion.

In the present paper we describe a genome-wide screen aimed at the identification of mutations affecting mouse development, in which we have chosen stage E10.5 for analysis. Although this is only one day later than in the screen of Kasarskis et al.,[10], we identified a comparatively quite different set of mutants.

We identified novel gene functions and isolated a series of novel mutant alleles for genes that were previously linked to mutant phenotypes. Three mutants isolated in this screen have been published elsewhere [12,13].

Results

Identified mutants

We analyzed the offspring of 150 G1 founders at embryonic day (E) 10.5 (Figure 1), a crucial stage for the development of the heart, limbs and neural tube. We identified at least 25 mutant lines characterized by a variety of reproducible phenotypes, including cardiac and nuchal edema, neural tube defects, a truncated anterior-posterior axis, situs inversus or the absence of limbs and lungs (Table 1; Figure 2). An exact number of mutant lines obtained is hard to give, as we terminated analysis of a number of potentially genuine mutant lines when mapping data were too confusing or reproducibility of the phenotype was doubtful. Therefore we can only give a minimum estimate. Amongst the mutants described included in Table 1 are several for which we have not yet identified the causative mutation.

Optimal ENU induction in mice is expected to lead to a mutation rate of approximately one mutation per 1.5 mega base pairs (Mb) [14]. Based on this mutation rate, a size of the mouse genome of approximately 2.5x10^9 Mb and assuming that approximately 1.5% of the genome encodes a protein, we anticipate that each G1 founder contains about 25 mutations that affect protein sequence.

A substitution mutation inDll1 leads to disturbed left-right patterning and axial truncation in 59459-2 mutants

A mutant characterized by a short tail as well as heart defects including situs inversus (Figure 3A), was mapped to Chromosome (Chr) 17 (5.5–27.5 Mb). Exons of genes in this interval were sequenced, which resulted in the discovery of an A to G transition (Figure 3B) in the gene encoding the Notch DSL ligand Delta-like1 (Dll1). This mutation, Dll1E26G, causes a glutamine-to-glycine substitution at amino acid position 26 (ENSMUSG00000014773). Glutamine 26 is highly conserved between species as well as in the Dll1-related protein Jagged/Serrate [13] and is positioned in the N-terminal domain 1 of the protein (Figure 3C).

Dll1 is known to be required for the formation and identity maintenance of the caudal somites [16] and for left-right asymmetry in the embryo [17]. Dll1 expression starts during the mid-streak stage in embryonic mesoderm. In the late-streak embryo expression is restricted to the posterior mesoderm, but it is absent from the node. At the 3-somite stage it is expressed in the presomitic mesoderm, the caudal halves of the condensed somites and in the neuroepithelium of the presumptive midbrain region [18].

Dll1E26G mutant embryos were posteriorly truncated (Figure 3A). Similar shortened anteroposterior axes have been previously reported for Dll1 mutants by Hrabe de Angelis et al. [19]. In addition the neural tube and the somites were irregularly shaped in the posterior part of mutant embryos (Figure 3H,I). Interestingly, an ectopic neural tube was occasionally present in Dll1E26G mutant embryos, ventrally of the primary neural tube and dorsally of the notochord (Figure 3F,G). Tbx6 mutant embryos often display an ectopic neural tube, which is accompanied by decreased levels of Dll1 expression in the mesoderm [20]. We are however not aware of similar reports for the Dll1 mutant itself. Finally, we observed randomized heart looping in Dll1E26G mutant embryos (Figure 3D,E) as previously noted by Przemeck et al. [17] for the previously characterized Dll1 mutant.

Notch signaling has been implicated in the development of the left-right axis, and heart looping and turning of the embryoare randomized in Dll1 mutants. This phenotype originates from the node, which displays morphological abnormalities in these mutants, including loss of monociliated cells [17]. The strong resemblance to the known knock-out null phenotype that the Dll1E26G embryos display indicates that the amino acid substitution observed is sufficient to abolish most or all of the biological function of the protein. A likely explanation is that Dll1E26 is essential in the binding of Dll1 to the Notch receptors, since the two N-terminal domains together with the DSL domain are responsible for the binding of DSL ligands to the Notch receptors [15,21,22]. The combination of mapping data with the similar phenotypes between Dll1E26G mutant embryos and the previously described Dll1 mutant makes allelism between 59459-2 and Dll1 very likely.

A premature stop codon in Ptprb/VE-PTP leads to early embryonic lethality in 59468-4 mutants

The 59468-4 mutant exhibits slight cardiac edema (Figure 4a–c). Genome-wide mapping linked it to a region on Chr 10 between 114 and 116.5 Mb. We noticed a striking resemblance to the
Table 1. Overview of mutants found in this screen (see Table S2 for additional information).

| Line*   | Phenotypeb | Chr | Mbpc | Affected gene | Mutation | Figure | Ref. |
|---------|------------|-----|------|---------------|----------|--------|------|
| S120-6B | Craniorachischisis | 15  | 71-83.2 | Scribble     |          |        |      |
| S120-6C | Cardiac edema         | 10  | 25-29  |              |          |        |      |
| S120-7  | Cardiac edema         | 17  | 78-84.5 | Ncx1         | N874K    |        | [13] |
| S120-8  | Open hindbrain        | 11  | 115-120 | Unknown      |          | 2B     |      |
| S9458-3 | Craniorachischisis    | 3   | 121.6-130.8 | Sec24b |          |        | [12] |
| S9459-2 | Situs inversus and short tail | 17  | 5.5-27.5 | Dll1         | E26G     | 3      |      |
| S9468-4 | Cardiac edema         | 10  | 114-116.5 | Ptprb       | Y693X    | 4      |      |
| S9622-3 | Cardiac edema         | 3   | 49-76   | Unknown      |          | 2E     |      |
| S97880-4 | NTD fore- and midbrain | 15  | 8-89.4  | PlexinB2     | E369G    | 5      |      |
| Amiko   | Growth arrest at E9.0 | 14  | 24-72   | Unknown      |          | 2L     |      |
| Cerbo   | Cardiac and nuchal edema | 2   | 165-166 | Unknown      |          | 2I     |      |
| Flanka  | Abnormal head, heart, NTD | 6   | 14.1-32.2 | Unknown |          |        |      |
| Koro    | Cardiac edema         | 11  | 3.2-17.6 | Unknown      |          | 2F     |      |
| Pootloos| No limbs              | 13  | -      | Fgf10        | L91P     | 6      |      |
| Linio   | Cardiac edema         | 11  | 94-98.7 | Unknown      |          | 2C     |      |
| Nevo    | Cardiac and nuchal edema | 8   | 77.4-98 | Unknown      |          | 2J     |      |
| Salsa   | Cardiac edema         | 6   | 67-71   | Unknown      |          | 2G     |      |
| Staartloos | Posterior truncation    | 11  | 55-66   | Wnt3a       | unknown  | 7      |      |
| Zoef    | Cardiac and nuchal edema | 19  | 33.5-33.8 | Unknown |          | 2H     |      |

*Trivial name of line.
**Major distinctive phenotype.
†Candidate interval after mapping.
doi:10.1371/journal.pone.0019357.t001

phenotype reported for mutants of one of the genes present in this interval, Ptprb, also known as E-PTP. Ptprb/VE-PTP mutants have previously been described as being defective in angiogenesis; vascular remodeling defects lead to a severely inflated pericardial sac and growth arrest at embryonic day 9.5 [23,24]. Baumer et al. also report that in their mutants the endocardium fails to attach to the myocardium, eventually leading to trabeculation defects; furthermore, they observed failure of intersomitic vessel development. These malformations strongly correlate with the expression pattern of Ptprb/VE-PTP mRNA, which is throughout development higher in the region on Chr 10 mentioned above, and sequencing of its coding regions revealed a T to A transversion at nucleotide position 445 of this 1998-amino acid long protein (Figure 4E). This early stop mutation causes a stop codon at the tyrosine at position 2079 (ENSMUST00000020363) (Figure 4D). This suggests that migration of insulin-like growth factor receptors into the cerebellar cytoarchitecture [27,28].

Exencephaly in the 59780-4 mutant is caused by a glutamic acid to glycine substitution in the Sema-domain of Plexin-B2

A mutant displaying exencephaly (Figure 5A–C) was mapped to chromosome 15 between 88.0 and 89.4 Mbp. Exons from the genes located in this segment were sequenced and the Plexin-B2 gene was found to contain an A to G transition only in DNA from mutants. This mutation predicts a glutamic acid to glycine substitution at amino acid position 369 E369G (Figure 5D,E). The membrane receptor Plexin-B2 is expressed in proliferating granule cell progenitors. Plexins are the receptors for Semaphorins; these ligands are involved in processes underlying proliferation, differentiation and migration in a variety of tissues [26].

Plexin-B2 mutants have been reported previously; they die at birth due to neural tube closure defects extending from fore- to hindbrain [27,28]. No molecular mechanism explaining this phenotype has been put forward. In the few mutants of the KO line that do not show a neural tube closure defect and survive until after birth, Plexin-B2 has been reported to have a role in maintaining the balance between differentiation and proliferation of the granule cells. Loss of this balance in Plexin-B2 mutants was said to result in over-proliferation of the differentiated granule cells that migrate into the cerebellum, leading to a severely altered cerebellar cytoarchitecture [27,28].

The amino acid that is changed in our Plexin-B2E369G mutant is localized in the Semaphorin-binding domain of the protein (Fig 5F), which may indicate that the glutamic acid at this position is essential for the binding of these Semaphorins to the cell surface receptor Plexin-B2. This possibility has not yet been tested. Given the combination of independent evidence, allelism of 59780-4 and Plexin-B2 is very likely, but remains formally unproven.

A leucine to proline substitution in FGF10 leads to absence of limbs and lungs in Pootloos mutants

The Pootloos mutant (Figure 6) was found on the basis of it lacking limb buds at E10.5 (Figure 6B,D). Alcian Blue and Alizarin Red
bone stainings at a later stage (Figure 6C,D) show the presence of the clavicle and anterior scapula. In the pelvic region, a rudimentary iliac bone was present but the fore- and hind limb bones are absent (Figure 6C,D). Transverse sections of E11.5 mutant embryos revealed the presence of primary lung buds, but no outgrowth and branching of the lungs (Figure 6E,F). This phenotype strikingly resembles the unique total loss-of-function phenotype of Fgf10 mutants described by Min et al. and Sekine et al. [29,30]. The mutants described by these authors are also characterized by the absence of limbs, retaining clavicle, anterior scapula and a rudimentary iliac bone. They have been reported to die shortly after birth due to the absence of the lungs [29,30]. After confirming linkage of the mutation to Fgf10 by limited mapping using a small number of SNPs, we proceeded by sequencing the Fgf10-coding region of Pootloos mutants and wildtypes. By exception, in this case the other candidates in this region were not sequenced. We identified a T to C transition predicted to cause a leucine to proline substitution at amino acid position 91 (ENSMUSP00000020363) (Figure 6H). This amino acid is part of a conserved domain common to all FGFs, but does not appear to be involved directly with functions characteristic of FGFs, like interactions with the FGF receptor or with heparin (see: http://www.ncbi.nlm.nih.gov/Structure/index.shtml; [31]; Figure 6G). The insertion of a proline is likely to have a major impact on essential aspects of the protein structure of the FGF-domain and may cause a total loss-of-function. The combination of independent evidence from a unique phenotype, the partial mapping and the presence of the mutation makes allelism of Fgf10 and Pootloos extremely likely.

Severe axial truncation in the Staartloos mutant

One mutant, named Staartloos, displaying a severe posterior truncation defect was found in the screen. Homozygous embryos had no tail, sometimes lacked hind limbs or had fused hind limbs (sirenomelia), and lacked caudal somites or vertebrae (Figure 7A,B). Mapping resulted in a candidate interval on chromosome 11 between 55 and 66 Mbp. All genes in this area were sequenced, but no mutations were found. The phenotype of this mutant line shows striking resemblances to the previously characterized knock-out phenotype of one of the genes present in the candidate interval. Wildtype mutants have also been described as posteriorly

Figure 2. Mutants identified in a recessive ENU-mutagenesis screen. A wildtype (A) embryo and the phenotypes of mutants indentified in the screen for which we thus far have not identified the causative mutation (B-L). doi:10.1371/journal.pone.0019357.g002
truncated, a complex of defects including disrupted development of notochord and tailbud and lack of caudal somites [32,33], based on failure of axial extension during early development. A striking phenotype previously reported for other Wnt3a mutant alleles is the occurrence of a partially duplicated neural tube. In work with Staartloos embryos, this phenotype has recently also been demonstrated (J. Deschamps, personal communication). Sequencing of a Wnt3a cDNA clone from a homozygous mutant, the coding region of genomic DNA, exon/intron transitions, 5′-untranslated regions neither revealed a mutation. We therefore crossed the Staartloos mutant with the Wnt3a KO mice [32] to explore non-complementarity of these two mutations. Double heterozygous mutant embryos were very similar to Staartloos mutants (Figure 7B,C), confirming that Staartloos carries a Wnt3a mutant allele. Possibly Wnt3a levels in the mutant are affected by a mutation in a regulatory element. Intriguingly, the causative mutation for the Wnt3a hypomorphic mutant Vestigial tail has never been reported either. In embryos homozygous for the Vestigial tail

Figure 3. A substitution mutation in Dll1 leads to situs inversus and posterior truncation. (A) Whole mount view of Dll1E26G and wildtype embryos. Note the shorter tail in the mutant. (B) An A-to-G mutation causing a glutamine-to-glycine substitution in the Dll1E26G mutant protein. (C) The mutation is localized in the N1 domain. SP, signal peptide; N1, N-terminal domain 1; N2, N-terminal domain 2; DSL, Delta-Serrate-Lag2 domain; EGF repeat, epidermal growth factor-like repeats; TMD, transmembrane domain; PDZL, PDZ (postsynaptic density 95, PSD-85; discs large, Dlg; zonula occludens-1, ZO-1) ligand motif. (D-I) H&E stainings on transverse sections at E9.5. The asterisk in E marks reversed looping compared to wildtype (D). NT, neural tube; Ht, heart. The arrow in G indicates an ectopic neural tube (F). Irregularly shaped neural tube (asterisks) and somites (arrows) in Dll1E26G mutants (H,I). Som, somites; Nc, notochord. doi:10.1371/journal.pone.0019357.g003

Figure 4. Cardiac edema in a mutant for Ptprb. Comparison of wildtype (A) and Ptprb mutant embryos (B,C). Note the cardiac edema in C. Sequence analysis shows a T to A mutation in Ptprb (D). Location of the PtprbY693X mutation (E). doi:10.1371/journal.pone.0019357.g004

Figure 5. A mutation in Plexin-B2 leads to exencephaly. Mutant embryos displaying closure defects throughout the brain region of the neural tube (A–C). Sequencing revealed an A-to-G point-mutation (D). Location of the Plexin-B2E369G mutation (E). doi:10.1371/journal.pone.0019357.g005
mutation, Wnt3a levels are reduced, suggesting that the mutation affects regulation of Wnt3a. Staartloos mutants contain more caudal vertebrae than Wnt3A KO embryos (Figure 7E-H), suggesting that the Staartloos mutation represents a hypomorphic condition for Wnt3a. Allelism of Staartloos with Wnt3a is therefore supported by four types of evidence: (i) genome-wide mapping; (ii) a phenotype very similar to that described in two mutant alleles of Wnt3a; (iii) a failure of the mutant to complement the Wnt3a phenotype. Therefore Staartloos is almost certainly allelic with Wnt3a.

Seven mutant lines carrying the same mutation

Seven mutants linked to an identical cardiac edema phenotype mapped in the same region of Chromosome 6 between 67 and 71 Mbp. Compound mutants of several combinations of these lines all led to the same phenotype, demonstrating that the same locus was affected. These lines must all have derived from a single spontaneous mutation present in some of the C57BL/6 mice used for theENU injections. We have combined these mutants under the name Salsa (Table 1 and Fig. 2) and are still in the progress of identifying the causative mutation.

Affected lung development in two PCP mutants

We identified two mutants in our screen that were characterized by the neural tube defect craniorachischisis in homozygous mutant embryos, Sec24bKrabbel and Scrib5120-6B; see [12]. In addition to the defects described above, we observed a lung development defect. At E17.5, the lungs of both mutants are smaller than those of wildtypes and the lobes are irregularly shaped (Figure 8A–F), with a clearly more severe phenotype seen in the Scrib mutant (Figure 8B,E; compare to Figure 8C,F). As we have shown that Sec24b is essential for adequate intracellular trafficking of Vangl2 in the neural tube [12], it is tempting to speculate that the lung

Figure 6. A leucine-to-proline substitution in the FGF domain of FGF10 leads to absence of limbs and lungs. Wildtype E10.5 embryo (A) and mutant embryo (B) displaying lack of limb buds at E10.5. In (A), yellow and orange arrows indicate location of fore and hind limbs (FL, HL), respectively, and similar arrows indicate in (B) the areas where they would have been expected to appear. (C,D) Comparison by bone staining of wildtype (C) and mutant (D) FGF10<sup>103P</sup> E15.5 embryos. In the mutant, clavicle (Clav), parts of the scapulae (Sc) and rudimentary iliac bones (Il) are present, but fore- and hind limb bones are absent. Sequencing revealed a T-to-C mutation (D). (E,F) Histological sections of E11.5 wildtype (E) and mutant (F) embryos at a thoracic level. ‘Lu’ in (F) indicates the lungs and arrows in G indicate the undeveloped lung buds in the mutant embryo. (G) Schematic representation of the alteration in the FGF protein, based on the detected mutation (H).

doi:10.1371/journal.pone.0019357.g006

Figure 7. Genetic interaction of Staartloos with Wnt3a. (A–D) Whole mount views of wildtype (A), Staartloos homozygous (B), Staartloos/Wnt3a transheterozygous (C) and Wnt3a KO mutant embryos (D). (E–H) Bone stainings of wildtype (E), Staartloos homozygous (F), Staartloos/Wnt3a double heterozygous (G) and Wnt3a KO mutants (H).

doi:10.1371/journal.pone.0019357.g007
Discussion

In this phenotype-based screen we have identified a series of new mutant alleles causing disrupted embryonic development. In some cases these new mutants represent unexpected genotype-phenotype associations while in other cases comparatively subtle mutations in previously studied genes reveal new findings at a molecular level.

The efficiency of our screen has been of similar efficiency compared to the comparable screens in the Anderson group (e.g., [3][10]). Kasarskis et al. [10] mention that 5–10% of founders tested showed a reproducible affected phenotype, and our data are in the same range. Exact meaningful numbers are difficult to give, since about 33 of the 150 founder lines were abandoned when we deemed the phenotype insufficiently reproducible, or when the initial mapping yielded unclear data; some of these lines may have represented real mutants. Of about 115 founders at least 4 G3 litters containing at least 30 embryos were analyzed.

We have no explanation for the fact that so few mutants identified by us display a defect linked to Sonic hedgehog signaling, as was the case in the screens mentioned above. Many of our mutants were identified by cardiac edema. This perhaps reflects the stage that we chose for the initial analysis of potentially mutant embryos, which is 1–2 days after the embryo becomes dependent on a functioning beating heart. In addition, cardiac edema is an endpoint of many different lethal defects.

Unexpected findings included the identification of a Scrib allele that must have arisen due to a ‘spontaneous’ (i.e. not ENU-induced) mutation of an FVB/N female, and the confusing findings with the Salsa mutant. In this case mutations were found that initially appeared to explain the phenotype; however the mutations were also present in some of the heterozygotes and must have been derived from genetic variation in the CB57/B6 mice used. This underscores the importance of checking the genetic purity of the mice strain used for the mutagenesis.

Figure 8. Affected lung development in the two PCP mutants obtained in our screen. (A–C) Dorsal and (D–E) ventral views of the lungs and hearts of (A,D) wildtype (B,E) Scrib and (C,F) Sec24b mutant embryos. Note the reduction in size of the mutant lung, in particular that of Scrib. (G–I) Sections of wildtype (G), Scrib mutant (H) and Sec24b mutant (I) lungs at E13.5. Ht and Lu (D) mark the heart and Lungs respectively, lumina (G) are labeled ‘Lum’. doi:10.1371/journal.pone.0019357.g008
PCP signaling and lung development

Scribble, Vangl2 and Sec24b have all been linked to the planar cell polarity (PCP) pathway that regulates the polarity in the plane of a sheet of epithelial cells and controls convergent extension of a tissue [12]; reviewed by [34]. The involvement of PCP signaling in lung development is supported by the following data: (i) In vitro lung branching is under the control of the downstream PCP factor RhoA [35]. (ii) Lung defects have been reported in mouse mutants for the ‘non-canonical’ Wnt5a gene [36]. Furthermore, over- and ectopic expression of Wnt5a in a cultured chick lung explant model confirmed its importance for lung development [37] and (iii) mutants for the PCP-related gene Fat6 [38] display branching defects in the kidney, an organ that shares similar signaling pathways with the lung and other organs containing branching tubes. These observations make it worthwhile further investigating the possible role of noncanonical Wnt signaling in lung development (see also: [39]). Recently Yates et al. [40] for the first time linked the core PCP pathway to lung development; they showed an abnormal size and shape of the lung lobes in Celsr1 and Vangl2 mutants. Both mutants have defects in lung branching and abnormalities of cytoskeletal architecture, as well as a reduced number and width of the airway lumina. The phenotype was confirmed in a lung explant model and a reduced response to FGF10 was demonstrated in this context. In vitro activation of Rho kinase restores lung branching in the Vangl2 and Celsr1 mutants. In addition to this, Celsr1 has a specific role in bifurcation of the lung buds. These authors also mention that they found a similar phenotype in their Scribble mutant [40]. We found, in addition to the small and irregular lung lobes, a disturbed cellular organization in the lumina in the Sec24b and Scribble mutants (Figure 3H,I), comparable to the double layers of epithelial cells in the Celsr1 and Vangl2 mutants [40]. Possibly, Sec24b and Scribble also direct lung development through the PCP pathway. The strong phenotype in the Scrib5120-6b mutant opens the possibility that there might be another crucial role for Scribble in lung development.

Conclusions

While reverse genetics (i.e., gene targeting technology) is powerful in that theoretically any desired genotype can be created, forward genetics is valuable as a parallel system and has a number of advantages: (i) the unbiased approach to identify genes of unknown function involved in development; (ii) the identification of single amino acid changes that alter protein function and (iii) the identification of novel phenotypes and functions of genes of known function that bypass lethality by the introduction of hypomorphic substitution mutations. A case in point for (i) is the Sec24b mutant, where we found the surprising link between Sec24b-dependent COPII-coated endoplasmic reticulum to Golgi protein transport, Vangl2 trafficking and PCP [12]. Examples of (ii) are the various mutants described in this paper in which a single missense mutation leads to a very strong phenotype, demonstrating in vivo the biochemical essentiality of the residue in question.

Recently, technological advances including ‘Next Generation Sequencing’ [41] have further reduced the efforts needed to isolate mutations. This will add greatly to the potential of ENU-mutagenesis screens, since the causative mutations can be found more rapidly. It should be possible to pinpoint the elusive mutations in Staatbosch and vestigial tail Wnt3a alleles in a relatively straight-forward manner, although this will also require analysis of reporter construct to test directly the mutations found.

Materials and Methods

Ethics statement

Animal experiments were conducted under the approval of the ‘dierexperimentencommissie’ (animal care committee) of the Royal Netherlands Academy of Arts and Sciences (KNAW), (permit numbers HL03.0204, HL04.0201, HL05.0201 and HL05.0203).

ENU-mutagenesis

40 C57BL/6 mice were injected three times intraperitoneally with 60–80 mg/kg bodyweight ethyl-N-nitrosourea (ENU) with one-week intervals. After a recovery period of 8–10 weeks the mice were crossed with FVB/N females to generate 150 G1 males. These ‘founder’ males were subsequently crossed with FVB/N to generate G2 females, who were crossed with their G1 father to generate G3 embryos. Each of these embryos is expected to be homozygous for 6.25% of the mutations initially induced in the C57BL/6 mice. These G3 litters were collected at E10.5 and screened for any perceivable malformations. DNA of the mutant and their littersmates was isolated for mapping. In rather numerous cases where a presumptive mutant was identified initially on the basis of severe cardiac edema at E10.5, we did further isolations at E9.5, which often allowed more relevant observations on the specific nature of the phenotype and also to avoid possible loss of embryos due to resorption. We screened a minimum of four G3 litters (of at least six, and on average more than eight embryos) produced by each G1 male. All mutant lines described in this paper are available from the authors to interested scientists. Lines were maintained by crossing against FVB/N until the mutation had been mapped. The exception was with the Scrib5120-6b line that arose from a mutation in a FVB/N mouse used for outbreeding [12].

DNA isolation

DNA was collected from embryos, yolk sacs (<E9.0) or ear clippings. The DNA was isolated by overnight incubation at 55°C while rotating in lysis buffer [TENS: 100 mMTris-HCl pH 8–8.5, 200 mMNaCl, 0.2% sodium dodecyl sulphate and 5 mMEthylendiaminetetraacetic acid (EDTA)] supplemented with 100 µg/ml proteinase K. From ear clippings hair was removed by centrifugation for 15 min at 12,000 x g. DNA was precipitated by adding 1 volume of 100% isopropanol at room temperature and vortexing for 5 min. To dissolve, the pellet was incubated with TE buffer (10 mMTris pH 7.5, 0.1 mM EDTA) overnight at 55°C while rotating. DNA was stored at 4°C.

Genetic mapping and genotyping

A panel of 192 SNPs discriminating between FVB/NJ and C57BL/6 DNA alleles was designed (see Table S1). These SNPs are spread equally over the genome and were used for initial genome-wide mapping using approximately 8 mutant and 8 wildtype or heterozygous DNAs isolated from each mutant line. Additional mapping using extra SNPs in the identified regions (http://phenome.jax.org/pub-cgi/phenome/mpdcgi?rtn=docs/home) reduced the candidate regions to intervals small enough to start sequencing coding regions (see Table 1). In a limited number of cases we gave priority to genes in this interval that had been previously characterized and shown to have phenotypes resembling our mutant. Mice and embryos were genotyped by PCR and sequencing of the mutation or SNPs in the identified interval. Generally between 70 and 100% of the genes in the candidate interval was sequenced, depending on the likeliness that found mutations corresponded to the phenotype observed. Table S2 summarizes mapping data for mutant lines discussed in this paper.

Sequence analysis

To amplify SNPs and exons of genes, a so-called touch-down program was used: 1’ at 94°C followed by 12 cycles of: [30°C at
92°C; 30" starting at 65°C and 0.6°C lower at each cycle; 30" at 70°C], followed by 20 cycles of: [30" at 92°C; 30" at 50°C; 30" at 72°C] followed by a final step of 3' at 72°C. GoTaq polymerase (Promega) was used according to the supplier’s instructions. Gene-specific primers were designed using software accessible at http://primer3.sourceforge.net and http://fimswill.nish.knaw.nl/PCR products were diluted and sequenced using BigDye v3.1 terminator (Applied Biosystems). The following protocol was used: 1' at 95°C, 30 cycles of 10" at 95°C, 5" at 50°C, 2" at 60°C. After the sequence reaction, the samples were purified using Sephadex G-50 Superfine (Sigma) or by ethanol/EDTA/Sodium Acetate precipitation. Sequencing was done at the Hubrecht institute using the standard RapidSeq protocol on 36 cm array. Mutations were identified using software based on Polyphred [42].

Miscellaneous procedures
Bone stainings were performed exactly as described by Beverdam et al. [43]. Hematoxylin and eosin staining were done according to standard procedures.

Supporting Information
Table S1 A 192-gene wide SNP panel. Listing of SNPs used in the genome-wide panel. First column, arbitrary number for identification; second column, rs numbers allowing retrieval from common databases (e.g. http://www.ncbi.nlm.nih.gov/projects/SNP/snp_blastByOrg.cgi); third column (‘Chr’), chromosome number; fourth and fifth columns provide expected nucleotide for C57Bl/6 and FVB/N, respectively.

Table S2 Mapping coordinates of mutant lines. Rs numbers correspond to the SNP database For each mutant the borders of the candidate region is given by the name (rs number) of the SNP [see http://www.ncbi.nlm.nih.gov/projects/SNP/snp_blastByOrg.cgi] as well as, in the last two columns, the chromosomal coordinates.

Acknowledgments
We thank Bart M. Smits for help in the initial setup of the screen and Shinji Takada for providing the Wnt5a KO mice.

Author Contributions
Conceived and designed the experiments: EC FM. Performed the experiments: CW LVG CK ER MV FM. Analyzed the data: CW HF ER VG EC FM. Contributed reagents/materials/analysis tools: VG EC. Wrote the paper: CW FM.
35. Moore KA, Pohc T, Huang S, Shi B, Alsberg E, et al. (2005) Control of basement membrane remodeling and epithelial branching morphogenesis in embryonic lung by Rho and cytoskeletal tension. Dev Dyn 232: 268–281.
36. Li C, Xiao J, Hormi K, Borok Z, Minoo P (2002) Wnt5a participates in distal lung morphogenesis. Dev Biol 249: 68–81.
37. Loscertales M, Mikels AJ, Hu JK, Donahoe PK, Roberts DJ (2008) Chick pulmonary Wnt5a directs airway and vascular tubulogenesis. Development 135: 1365–1376.
38. Saburi S, Hester I, Fischer E, Pontoglio M, Eremina V, et al. (2008) Loss of Fat4 disrupts PCP signaling and oriented cell division and leads to cystic kidney disease. Nat Genet 40: 1010–1015.
39. Morrisey EE, Hogan BL (2010) Preparing for the first breath: genetic and cellular mechanisms in lung development. Dev Cell 18: 8–23.
40. Yates LL, Schnatwinkel G, Murdoch JN, Bogani D, Formstone CJ, et al. (2010) The PCP genes Celsr1 and Vangl2 are required for normal lung branching morphogenesis. Hum Mol Genet 19: 2251–2267.
41. Metzker ML (2010) Sequencing technologies - the next generation. Nat Rev Genet 11: 31–46.
42. Nickerson DA, Tobe VO, Taylor SL (1997) PolyPhred: automating the detection and genotyping of single nucleotide substitutions using fluorescence-based resequencing. Nucleic Acids Res 25: 2745–2751.
43. Beverdam A, Brouwer A, Reijnen M, Korving J, Meijlink F (2001) Severe nasal clefting and abnormal embryonic apoptosis in Alx1/Alx4 double mutant mice. Development 128: 3975–3986.