Development and regeneration of the nervous system requires the precise formation of axons and dendrites. Kinases and phosphatases are pervasive regulators of cellular function and have been implicated in controlling axodendritic development and regeneration. We undertook a gain-of-function analysis to determine the functions of kinases and phosphatases in the regulation of neuron morphology. Over 300 kinases and 124 esterases and phosphatases were studied by high-content analysis of rat hippocampal neurons. Proteins previously implicated in neurite growth, such as ERK1, GSK3, EphA8, FGFR, PI3K, PKC, p38, and PP1a, were confirmed to have effects in our functional assays. We also identified novel positive and negative neurite growth regulators. These include neuronal-developmental regulated kinases such as the activin receptor, interferon regulatory factor 6 (IRF6) and neural leucine-rich repeat 1 (LRRN1). The protein kinase N2 (PKN2) and choline kinase α (CHKA) kinases, and the phosphatases PPEF2 and SMPD1, have little or no established functions in neuronal function, but were sufficient to promote neurite growth. In addition, pathway analysis revealed that members of signaling pathways involved in cancer progression and axis formation enhanced neurite outgrowth, whereas cytokine-related pathways significantly inhibited neurite formation.

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Keywords: bioinformatics; development; functional genomics; metabolic and regulatory networks; neuroscience

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Introduction

In neural development, neuronal precursors differentiate, migrate, extend long axons and dendrites, and finally establish connections with their targets. The initial steps of neurite formation and the subsequent specification of axons and dendrites have been widely studied. Clinical conditions such as spinal cord injury, traumatic brain injury, stroke, multiple sclerosis, Parkinson’s disease, Huntington’s disease, and Alzheimer’s disease are often associated with a loss of axon and/or dendrite connectivity and treatment strategies would be enhanced by new therapies targeting cell intrinsic mechanisms of axon elongation and regeneration.

Phosphorylation controls most cellular processes, including the cell cycle (Vandenheuvel and Harlow, 1993), proliferation (Behrens et al., 1999), metabolism (Nimmo and Cohen, 1978), and apoptosis (Xia et al., 1995). Neuronal differentiation, including axon formation and elongation, is also regulated by a wide range of kinases and phosphatases (reviewed in Arimura and Kaibuchi, 2007). For example, the non-receptor tyrosine kinase Src (Entrez symbol: SRC) is required for cell adhesion molecule-dependent neurite outgrowth (Ignelzi et al., 1994) and the phosphatase Calcineurin (PPP3CA) is required downstream of calcium waves to transiently inhibit the extension of Xenopus spinal neurons (Lautermilch and Spitzer, 2000). In addition to individual kinases and phosphatases, signaling pathways like the MAPK, growth factor signaling, PIP3, cytoskeletal, and calcium-dependent pathways have been shown to impinge on or control neuronal process development (Lazarovici et al., 1987; Kuo et al., 1997; Morooka and Nishida, 1998; Wu and Cline, 1998; Huang and Reichardt, 2003; Menager et al., 2004; Chen et al., 2006).

Recent results have implicated GSK3 (Dill et al., 2008) and PTEN (Park et al., 2008) as therapeutically relevant targets in axonal regeneration after injury. However, these and other experiments have studied only a small fraction of the total kinases and phosphatases in the genome. Because of recent advances in genomic knowledge, large-scale cDNA production,
and high-throughput phenotypic analysis, it is now possible to take a more comprehensive approach to understanding the functions of kinases and phosphatases in neurons. We performed a large, unbiased set of experiments to answer the question ‘what effect does the overexpression of genes encoding kinases, phosphatases, and related proteins have on neuronal morphology?’ We used a ‘high-content analysis’ approach to obtain detailed results about the specific phenotypes of neurons. We chose to study embryonic rat hippocampal neurons because of their stereotypical development in vitro (Dotti et al., 1988) and their widespread use in studies of neuronal differentiation and signaling. We transfected over 700 clones encoding kinases and phosphatases into hippocampal neurons and analyzed the resulting changes in neuronal morphology. Many known genes, including PP1a, ERK1, p38, ErbB2, atypical PKC, Calcineurin, CaMK2, FES, IGFR1, FGFR, GSK3, PDK1, PIK3, and EphA8, were observed to have significant effects on neurite outgrowth in our system, consistent with earlier findings in the literature. Importantly, we also identified a number of genes not previously known to affect process growth. Combining the morphological data with information about protein sequence and molecular pathways allowed us to connect families of related proteins with novel functions in neurite development, and to implicate some signaling pathways in the regulation of neurite growth for the first time. Overall, our results provide a more complete picture of the kinases and phosphatases regulating neuronal growth, and suggest a number of testable hypotheses regarding the signaling pathways involved.

Results

A large-scale gain-of-function analysis in primary mammalian neurons

Electroporation-mediated transfection was used to overexpress kinases and phosphatases in embryonic rat hippocampal neurons. These neurons quickly adhere to laminin-coated plates, initiating neurite growth within hours (Esch et al., 1999). By 48 h, neurons typically possess several minor neurites and one major neurite (likely to develop into the axon) (Dotti et al., 1988). We marked transfected neurons by cotransfection with mCherry, a red fluorescent protein (RFP) (Shaner et al., 2004); transfection efficiency averaged 17.3% (95% confidence interval (95 CI), 16.6–18%) of the BII-tubulin-positive neurons. Only transfected neurons were analyzed; neurons were defined as transfected (RFP+; Figure 1B and D, arrowheads) if their RFP intensities were greater than 2 s.d. above the mean of non-transfected controls (Figure 1E and F). Control experiments demonstrated that >80% of RFP+ neurons were cotransfected with the gene of interest (data not shown). Except when measuring the percent of neurons with neurites (%Neurite+), we considered neurons for further analysis only if they had at least one neurite >10 μm (Neurite+; Figure 1A and B) to avoid measuring potentially non-viable neurons (Figure 1C and D).

We obtained quantitative data for many cellular and neuronal morphological parameters from each neuron imaged. These included nuclear morphology (nuclear area and Hoechst dye intensity), soma morphology (tubulin intensity, area, and shape), and numerous parameters of neurite morphology (e.g. tubulin intensity along the neurites, number of primary neurites, neurite length, number of branches, distance from the cell body to the branches, number of crossing points, width and area of the neurites, and longest neurite; Supplementary Figure 1). Other parameters were reported on a ‘per well’ basis, including the percentage of transfected neurons in a condition (%RFP+), as well as the percentage of neurons initiating neurite growth (%Neurite+). Data for each treatment were normalized to the control (pSport CAT) within the same experiment, then aggregated across replicate experiments.

Validation of normalization and use of transfected neurons

We constructed a linear model incorporating experimental and treatment terms to test the validity of our normalization and our method for selecting transfected neurons (Figure 2C). The main sources of variance were the treatment (overexpression of kinases and phosphatases) and various aspects of experimental technique (animals, cells, time of prep, transfection, etc).

\[ NTL_{ij} = C + E_i + T_j + ET_{ij} + \epsilon_{ij} \]

where C=constant, E=experiment effects, T=treatment effects, and \( \epsilon = \) error.

The linear models fit the data appropriately, with coefficients of RFP= 0.62, 0.59 and RFP+ 0.87, 0.84 (for raw and normalized data models, respectively). The models were analyzed to determine the respective contributions of the treatment term (representing changes specifically because of overexpression), and the experimental term (representing noise). When analyzing cells defined as non-transfected, the experimental (noise) contribution predominated, both for raw and normalized data (Figure 2C). However, for cells defined as transfected (RFP+), the treatment effect increased (to 34%) and the noise contribution decreased (to 32%), even for raw data. For normalized data from transfected cells, the treatment contribution increased to 51%, whereas the experimental contribution became negligible (6%). We concluded that it was appropriate to analyze the RFP+ neurons, and that normalization significantly reduced the noise in the screen, revealing the effect of cDNA overexpression. Notably, the interaction term maintained a sizable contribution in our model (not shown), suggesting that a non-linear model would be able to explain more of the variance than our simple linear one.

Neurite number is regulated separately from neurite elongation

Correlations among the 19 normalized parameters were analyzed for neurons transfected with all kinase and phosphatase clones (Figure 2A). Primarily, this approach was taken to simplify the number of variables. Five groups of parameters as well as four single parameters emerged from the correlation analysis. Parameter groups comprised clusters of red intensity measures, soma geometry measures, neurite crossing, branching and neurite area, and neurite length measures (defined in
Supplementary Figure 1). On the basis of this analysis, the primary variables that define the neurite morphology are primary neurite count, neurite average length, and average branches. Interestingly, primary neurite count was not well correlated with neurite length or branching (Figure 2A). The Pearson correlation coefficient \((r^2)\) between the number of primary neurites and the average length of the neurites was 0.332 (95 CI, 0.247–0.412), and between the number of primary neurites and average branching was 0.227 (95 CI, 0.146–0.302). In contrast, the correlation coefficient of average...
branching with neurite average length was 0.670 (95 CI, 0.626–0.712) (Figure 2B). The primary neurite count was well correlated with neurite total length (NTL). This is expected as NTL is defined as (average length × primary neurite count) (Supplementary Figure 1).

There are at least two possible explanations for the low correspondence between primary neurite number and either neurite length or branch number. First, the number of primary neurites in embryonic hippocampal neurons could be very stable and imperturbable. Indeed, primary neurite number did not vary a great deal across the entire data set (2.13 ± 1.14 s.d.). However, we identified a group of genes (see below) that significantly perturbed primary neurite count, rendering this explanation unlikely. Therefore, a more likely possibility is that signaling mechanisms underlying the neurite number determination are different than those controlling length/branching of the neurites. A mechanism for this phenomenon has recently been examined (Shelly et al., 2010).

Phenotypic perturbation after overexpression of kinases and phosphatases

Neurons were transfected with each of 724 cDNA plasmids, from the NIH Mammalian Genome Collection (MGC) (Gerhard et al., 2004), mapping onto 622 different human and mouse genes. As 130 of these genes were orthologs, 492 unique genes were screened. After image/tracing validation, a number of clones were removed from the analysis (because of poor tracing or low cell number), leaving 449 unique genes (plus five control genes) in the final analysis.

To identify relevant functional classes of neuronal phenotypes, three key parameters were considered—primary neurite count, average length, and branching. Changes in these parameters resulting from gene overexpression were examined using hierarchical clustering (Figure 3). Of the 454 unique genes (including controls), 59 had statistically significant positive effects in neurite count and length, and are grouped on the left side of the heat map in green. Only nine genes increased both the primary neurite count and the average neurite length, whereas 23 genes increased neurite average length. An additional 32 genes increased the primary neurite count, but with little or no increase in length. Only a few of the genes assayed were significant determinants of both neurite length and neurite count. It is likely that these genes control general neurite initiation, whereas the other phenotypic classes regulate more specific cellular programs. A majority of genes with significant effects (92) were inhibitory to neurite growth.

We identified several genes that significantly altered neurite growth. For example, choline kinase α (CHKA) promoted positive growth enhancement for all parameters after over-
expression in hippocampal neurons (Figure 3F). Origin recognition complex subunit 4-like and the TGF-β receptor 2, increased the length of hippocampal neuron processes (Figure 3G and H). Origin recognition complex was previously reported to regulate dendrite development (Huang et al., 2005). Inositol polyphosphate phosphatase (INPP5B) overexpression increased the number of primary neurites and led to decreased neurite length (Figure 3J). Although these genes had enhancing effects on laminin, they had little effect on neurite growth on the artificial substrate polylysine (Supplementary Figure 2). The strongest inhibitory genes produced very short neurites with only a few neurons initiating growth (e.g. atypical PKC iota, PRKCI; Figure 3K). These inhibitory genes did not generally affect survival (Supplementary Figure 3). Overall, we identified several neurite growth activators, and many neurite growth inhibitors in hippocampal neurons. The full table of results is provided in Supplementary Table 1.

Effects of known neurite-regulatory genes on neuronal morphology

We selected a group of genes that were tested in our experiments and had been previously implicated in neuron development, neuritogenesis, axon elongation, or neuronal polarity. Table I reports the results of overexpression of these genes in hippocampal neurons for four parameters: nuclear intensity (increases reflect nuclear condensation; Marcus et al., 1979), primary neurite count, neurite average length, and average branching. The latter three parameters were chosen because they defined the three distinct groups of neurite variables. Genes that had significant effects (after correction for multiple comparisons) are indicated. As expected, the protein kinase A catalytic subunit (PRKACA) and the atypical PKCs (PRKCI, PRKCH) strongly perturbed the neuron’s phenotype. Overexpression of these kinases was sufficient to inhibit neurite growth. PKA activity has previously been suggested to affect growth on laminin negatively (Bixby, 1989), but also has been linked to positive regulation of neurite growth (Kao et al., 2002). It is possible that our (inhibitory) results with PKA reflect mislocalization because of expression of the catalytic subunit in the absence of the regulatory subunit. Note that although overexpression of PKA inhibited neurite length, it did not increase the nuclear intensity, suggesting cell death was not the cause (Table I). To determine whether PKA’s effect was due to its kinase activity, we compared expression of wild-type PRKACA to that of an inactive mutant. Transfection of PRKACA led to strong inhibition of neurite growth (Figure 4B), whereas transfection of a kinase-dead mutant (K73A; Figure 4C) did not alter neuronal morphology (compare Figure 4A). Therefore, PRKACA’s kinase activity was required for its inhibition of neurite growth.

Neuronal morphology is altered by expression of members of kinase and phosphatase families

Related proteins are often involved in similar neuronal functions. For example, families of receptor protein tyrosine phosphatases are involved in motor axon extension and guidance in both Drosophila and in vertebrates (Desai et al., 1997; Stepanek et al., 2005), and a large family of tyrosine kinases regulates guidance of retinotectal projections, motor axons, and axons in the corpus callosum (Brennan et al., 1997; Klein, 2001; Mendes et al., 2006). We therefore asked whether families of related genes produced similar phenotypes when overexpressed in hippocampal neurons. Our set of genes covered 40% of the known protein kinases (Supplementary Table 2), and many of the non-protein kinases and phosphatases.

Genes were sorted into one of five categories before the sequence alignment to produce distinct trees: protein kinases (Figure 5A), non-protein kinases (Figure 5B), protein phosphatases (Figure 5C), other phosphatases (Figure 5D), and other genes (Figure 5E). Phylogenetic trees were used to determine amino-acid similarity and infer evolutionary distance. Related families were thus clustered into branches and subbranches. Genes identified as protein kinases (Manning et al., 2002; Caenepeel et al., 2004) were aligned using the amino-acid sequences of the kinase domains only, whereas other genes were aligned by their entire sequences. The alignment successfully clustered the genes into known families. For example, the protein kinase tree (Figure 5A) matched the organization of previously identified protein kinase groups (Manning et al., 2002; Caenepeel et al., 2004). Also, 81% of the genes with obvious siblings (i.e. PPP2CA, PPP2CB) were linked to their sibling in the trees.

To establish the neuronal phenotype for individual genes within families, we plotted the functional data for NTI on markers overlying the trees (Figure 5). The neuronal response observed after the overexpression of individual kinases and phosphatases was diverse across gene families. Expression of most kinase and phosphatase genes reduced the length of the neurites compared with controls (Figure 5, red markers). Some families were particularly inhibitory. For example, BMP/actin-kinase receptors, PKAs/PKCs (PRKACx, PRKCx), and most protein serine/threonine phosphatases (PPP branch) inhibited neurite extension.

Gene families commonly exhibit redundant function (Urrutia et al., 1997; Xian and Zhou, 2004). Redundant gene function has often been identified when two or more knockouts are required to produce a phenotype (Heber et al., 2000). Our technique allowed us to measure whether different members of gene families had similar (potentially redundant) or distinct effects on neuronal phenotype.

To determine whether groups of related genes affect neuronal morphology in similar ways, we used sequence alignment information to construct gene clusters (Figure 6A). Genes were clustered at nine different thresholds of similarity (called ‘tiers’). The functional effect for a particular parameter was then averaged within each cluster of a given tier, and non-parametric statistics were performed to determine the significance of the effect. Results from the neurite initiation parameter are shown in Figure 6B. Here, the giant cluster containing every gene has an average near control level and is colored white (Figure 6B, bottom tier, labeled 1 on the left). As the threshold is increased, clusters of fewer but more closely related genes are constructed, and their mean effect on the phenotype is displayed by the color of the rectangles in the heat map (red=decrease, green=increase, diamonds and asterisks indicate significance).
In all, 48 kinases and phosphatases previously reported for their involvement in neural process development and polarity. Columns of the table refer to intensity of the key neurite parameters analyzed. This family of kinases consisted of a variety of enzymes, mostly sugar and lipid kinases, with the strongest effects coming from IRF6 and CHKA.

### Molecular pathways revealed by neuronal phenotypes

Over the past 10–20 years, literally hundreds of neuronal growth and regeneration-associated genes have been identified (e.g. Table I). However, determining which of these genes

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**Table I  Known neural growth regulators**

| Common name | Symbol | Note | Citations |
|-------------|--------|------|-----------|
| AKT2        | AKT2   | Inhibition correlated with decrease | Namikawa et al (2000) |
| CAMK1d      | CAMK1D | Inhibiting decreased length | Wayman et al (2004) |
| CAMK1g      | CAMK1G |                |           |
| CAMK2a      | CAMK2A | Slowed dendritic growth, mediates GABA induced neurite growth | Wu and Cline (1998); Borodinsky et al (2003) |
| ERK2        | ERK2   | Activated form induces outgrowth | Kolkova et al (2000); Robinson et al (1998) |
| Epha4       | EPHA4  | Better regeneration in Epha4 KO | Goldshmit et al (2004) |
| EphA7       | EPHA7  | Better recovery after EphA7 reduction | Figueroa et al (2006) |
| ErbB2       | ERBB2  | Neuregulin promotes through erbB2 | Bermingham-McDonogh et al (1996) |
| FAK         | FAK    | GF + Integrin > FAK > PC12 growth | Ivankovic-Dikic et al (2000) |
| FGF receptor1 | FGFR1 | Overexpression-promoted outgrowth | Hausott et al (2008); Lin et al (1996) |
| FGFR1       | FGFR1  |                     |           |
| FGFR4       | FGFR4  |                     |           |
| Fyn related (GTK) | FYN | Fyn-mice lack NCAM neurite growth | Beggs et al (1994) |
| GSK3A       | GSK3A  | Has distinct function from below | Yoshimura et al (2005); Lee et al (2007) |
| GSK3B       | GSK3B  | Inhibition of GSK3b promotes | Yoshimura et al (2005) |
| IGFR1       | IGFR1  | Receptor is essential for Hipp polarity | Sosa et al (2006) |
| ILK         | ILK    | Overexpress Wt ILK stimulates growth | Ishii et al (2001) |
| JNK2        | MAPK9  | JNK required for axon formation | Oliva et al (2006) |
| JNK3        | MAPK10 |                     |           |
| MARK2       | MARK2  | Ectopic expressed led to loss of axons | Chen et al (2006) |
| MEK3        | MAPK3  | MEK required for TrkB growth | Atwal et al (2000) |
| MEK5        | MAPK5  |                     |           |
| ORC4L       | ORC4L  | Knockdown-reduced branching | Huang et al (2005) |
| PAK4        | PAK4   | Membrane targeting increase growth | Daniels et al (1998) |
| PI3K        | PIK3   | Activated by PI3K | Alessi et al (1997) |
| PKA catalytic A | PRKACA | Inhibit PKA stops forskolin growth | Kao et al (2002); Chijiwa et al (1990) |
| PKA catalytic B | PRKACB |                     |           |
| PKC eta     | PRKCH  | Cell polarity and Par complex | Chen et al (2006); Lin et al (2000) |
| PKC iota    | PRKCI  |                     |           |
| Raf1        | RAF1   | Activated raft–axon elongation | Markus et al (2002) |
| SRC         | SRC    | Required for polarization | Kishi et al (2005) |
| SRC KO      | SRC    | SRC KO impairs growth | Ignezi et al (1994) |
| CDC42       | CDC42  | Inhibits PKA | Nikolic et al (1998) |
| p38 a       | MAPK14 | Required for PC12 outgrowth; inhibiting enhances growth | Morooka and Nishida (1998); Myers et al (2003) |
| p38 b       | MAPK11 |                     |           |
| p38 d       | MAPK13 |                     |           |
| p38 g       | MAPK12 |                     |           |
| Calcineurin | PPP3CA | Ca ++ waves inhibit though Calcin | Lautermilch and Spitzer (2000) |
| PP1a        | PPP1CA | Dephosphorylates PKA | Tang et al (2003) |
| PTEN        | PTEN   | PTEN KO retinal regrowth after crush | Park et al (2008) |
| PTPRα       | PTPRA  | Drosophila | Desai et al (1997) |
| SHP2        | PTPN11 | NGF growth inhibited by mutant Shp2 | Chen et al (2002) |

In all, 48 kinases and phosphatases previously reported for their involvement in neural process development and polarity. Columns of the table refer to intensity of the Hoechst nuclear dye (1), primary neurite count (2), neurite average length (3), and average branching (4). Effect for each parameter (+, ++, +++; −, −−, −−−; P<0.01, corrected by Benjamini–Hochberg), compared with controls. Genes are identified by their common name, official Entrez symbol, and official full name. Genes are arranged with the kinases on the top, and the phosphatases below.
are primary regulators of process development has been challenging. It is possible to refine the understanding of these genes with existing pathway information. One hurdle is that most genes with pathway annotations are members of multiple pathways. Of the 550 genes we successfully screened, 46% had pathway annotations, and of those, 66% were members of more than one pathway. Of the latter, half were members of four or more pathways. We performed pathway analysis for 79 different pathways in 24 categories using Kyoto Encyclopedia of Genes and Genomes (KEGG) (Kanehisa et al., 2004) (http://www.genome.jp/kegg/pathway.html) (Figure 7A and B). This analysis was displayed in the form of a heat map (Figure 7C). KEGG provided the classification of main categories (Figure 7A), subcategories (Figure 7E), and the pathways (Figure 7B). Genes were then placed in these pathway ‘bins.’ The color of the bin indicates the strength and the direction of the effect observed (Figure 7C and D). Pathways involved in proliferation of specific cancers, dorso-ventral axis formation, cytokine signaling, and ErbB signaling each had genes that were significant in their ability to modify neuronal morpho-

Figure 4  Overexpression of PKA catalytic subunit α inhibits neurite growth. Its kinase activity is responsible for the phenotype change. (A–C) Hippocampal neurons transfected with control (mCherry, A), PKA catalytic subunit α (PRKACA, B), and kinase-dead mutant of PKA (PRKACA K73A, C). Arrowheads indicate PRKACA-transfected neurons; arrows indicate neurons expressing the kinase-dead mutant of PKA. The neurons’ processes are visualized by βIII-tubulin (green), and mCherry coexpression (red). Scale bar=100 μm.

Figure 5  Families of kinases and phosphatases modify neurite outgrowth. Screened genes were aligned by sequence then displayed as phylogenetic trees, representing evolutionarily related proteins. Genes were separated into five primary categories, protein kinases (A), non-protein kinases (B), protein phosphatases (C), non-protein phosphatases (D), and others (E). Protein kinases were further subdivided into seven families (based on kinase domain sequences). Branch lengths on the trees represent the amino-acid change from parent to sibling and therefore the evolutionary ‘distance’ between genes. Labels beside the branches are the official Entrez Gene symbol. Circular markers were overlaid atop the trees, with the color of the marker indicating either no change in neurite length from control (white), an increase (green) or decrease (red). KLF4 and KLF7 are transcription factors that are regulators of neurite growth (Moore et al., 2009; Blackmore et al., 2010), and were included in the screen as additional controls. Source data are available for this figure at www.nature.com/msb.
Figure 6  Bioinformatics identify putative families of active genes. (A) Genes overexpressed in hippocampal neurons, arranged by phylogenetic relationships (as in Figure 5, but ‘unrolled’) where the most closely related genes are next to each other. The five trees are preserved in the five sections (columns) of the graph. (B) Cluster heat map for the frequency of neurite initiation. At the bottom (labeled 1), all of the genes are combined into one large cluster, where there was no net effect on neurite growth (white). Going up from the bottom, genes are grouped in smaller, more focused families, with 3, 9, 21, 45, 88, 199, and 347 families per tier (row of the heat map). Each rectangle represents a family of genes (corresponding to the labels in A) that has a mean effect that reduced (red), or increased (green) neurite initiation. Non-parametric statistics were performed on each family, where diamonds indicate values for spiked bootstrap (**P < 0.01). Asterisks indicate significance corrected by Benjamini–Hochberg (*P < 0.05, **P < 0.01). (C) Compilation of significant families for four parameters: primary neurite count, frequency of neurite initiation, neurite average length, and average branching. Colored rectangles demarcate the extent of the family (for instance CDKN3, PTEN, PTAP1, CDC14A&B), the color the direction and magnitude of the effect, with labels giving the common names for the genes. Note that the large family of non-protein kinases (from FUK to PFKL) has smaller nested families with even greater effects. The calcineurins (PPP3CA&B), and the kinases IRF6, CHK4, and CDK5R1, had the greatest ability to perturb neurite growth. Source data are available for this figure at www.nature.com/msb.
Pathways provide a conceptual framework to understand molecular dynamics in cells. We analyzed hundreds of genes for functional relationships, and found that several pathways implicated in cancer progression and dorsal-ventral pattern formation contain kinases and phosphatases with substantial abilities to alter neurite morphology.

**Novel growth-associated genes**

Our study uncovered a number of genes, including developmentally active genes, not previously linked to the regulation of axon/dendrite growth. For example, activin receptor 1 (ACVR1) is involved in Xenopus axis formation (Hemmati-Brivanlou and Melton, 1992) and downregulated during peripheral nerve regeneration (Morita et al., 1996). Both ACVR1 and ACVR1L (and their isoforms) reduced neurite growth in our assays. IRF6 is also developmentally regulated during early development (Hatada et al., 1997) and its expression potentiated neurite growth in hippocampal neurons. CHKA, which strongly increased hippocampal neurite growth and neurite initiation, is an essential gene (Wu et al., 2008) that has been observed in nerve endings (synaptosomes) (Spanner and Ansell, 1979). Elucidating the mechanism through which CHKA influences neurite growth will be a fruitful future direction of study. SBK1 (SH3-binding kinase 1) is a novel and mostly uncharacterized serine/threonine kinase that is highly expressed in the brain (Nara et al., 2001). Two mouse clones of this gene with different UTRs inhibited neurite growth in our assays. Another serine/threonine kinase, microtubule-associated serine/threonine kinase-like (MASTL), which inhibited neurite outgrowth, is known to interact with microtubules. A non-protein kinase, neural leucine-rich repeat 1 (LRRN1), is highly expressed in early somitic myoblasts (Haines et al., 2005) and inhibited neurite growth when overexpressed.

Neurite growth activators were very rare in the AGC arm of the protein kinase tree, with the exception of protein kinase N2 (PKN2), known to be activated by Rho GTP, and regulate cell cycle proteins such as Cdc25B (Schmidt et al., 2007). Another AGC kinase, ribosomal protein S6 kinase/mitogen and stress activated kinase/p90rsk (RPS6KA4), inhibited neurite growth. RPS6KA4 has been shown to be required for CREB and ATF phosphorylation in fibroblasts (Wiggin et al., 2002), as well as being involved in cAMP and insulin-related proliferation (Coulomval et al., 2000). It has previously been implicated in regulating CAM-mediated neurite growth (Wong et al., 1996).
Two protein phosphatases, both members of the myotubulinarin family, significantly decreased neurite growth. Myotubulinarin related 2 (MTMR2) is known to be involved in myelin neuropathies, but is currently thought to act in Schwann cells, not neurons (Bolis et al., 2005). SET-binding factor 1 (SBF1) a pseudo-protein phosphatase, has been seen to decrease proliferation after its forced expression in fibroblasts (Firestein and Cleary, 2001). We observed SBF1 to decrease neurite growth after overexpression in hippocampal neurons.

Other genes with strong effects have either less known about them, or no obvious connection to neural function or pathways. The serine threonine kinase protein serine kinase H1 is involved in Golgi maintenance (Brede et al., 2003) and its overexpression enhanced growth. CMPK1 (cytidine monophosphate UMP-CMP kinase 1) a kinase involved in pyrimidine synthesis, inhibited neurite growth when overexpressed. The protein phosphatase protein phosphatase 1K was inhibitory, whereas protein phosphatase EF hand calcium-binding domain 2 (PPEF2) and sphingomyelin phosphodiesterase 1 acid lysosomal (SMPD1) were strong growth promoters in our screen, but the underlying mechanisms are unclear.

Candidates for future study from genetic and pathway analyses

Our phylogenetic analysis identified several families, such as the chemokine ligands, protein tyrosine phosphatases, and a large group of non-protein kinases, with members that strongly perturbed neurite outgrowth. Several of these were expected, especially the Calcineurins, Cdc14s, PI3K (Da Silva et al., 2005), and atypical PKCs. Other families were not expected to affect neurite growth, but did. These include a group of sugar kinases, NADK, and others. Not every member of this family had the same effect, but these enzymes generally potentiated neurite growth over control (surprising considering the neurons were growing on laminin, which strongly promotes neurite growth in the control state (Baronvanevercooren et al., 1982)). These non-protein kinases will be interesting to study to determine their functions in neuronal process development.

Pathway analysis revealed that genes in several cancer pathways were highly active. This is not unexpected, as growth of neurons is tied to the same cascades that lead to proliferation of cancer cells (Nakagawara, 2001). These pathways dominate the analysis, either because many of their genes were selected for study, and have been previously studied, or that these signals (as activated by overexpression) are quite strong compared with other pathways. Finally, the signal transduction pathways ‘neuroactive ligand–receptor interaction’ and ‘cytokine–cytokine receptor interaction’ have members that produced strong effects in opposite directions (Supplementary Figure 4, purple blocks). The study of these complete pathways will be an important undertaking for future analysis.

Comparison with other screens

Recently, three groups have used RNAi to test the necessity of various genes on neurite development or cell migration, which is likely to involve overlapping mechanisms (Enomoto et al., 2001; Maness and Schachner, 2007). Loh et al. (2008) used the SH-SY5Y cell line in an siRNA screen of 750 kinases, many of which were also assayed in our experiments. Over half of the ‘hit’ siRNAs targeted genes in the tyrosine kinase or tyrosine kinase-like families. The intersection of hit genes from Loh et al. and genes from our study is listed in Supplementary Table 3. ERBB2, RYK, PRKAB1 enhanced neurite growth after overexpression in our assays and reduced neurite length after knockdown in the Loh screen. Conversely, PRKCI, PRK1, PDPK1 NEK3, MASTL inhibited neurite growth after overexpression in our assays and enhanced axon growth after knockdown in the Loh screen. Overexpression/knockdown of other genes (CDK9, FES, STK38L) led to the same phenotype in both screens. An siRNA screen by Sepp et al. (2008) used Drosophila primary neurons to study the disruption of neural phenotypes in a genome-wide screen. They found that siRNAs for over 100 genes significantly perturbed neuronal phenotypes, leading to reduced elongation, excessive branching, loss of fasciculation, or blebbing (Sepp et al., 2008). The orthologs of four of the active genes from Sepp et al were also screened in our experiments. Two of these, ZAP70 and HTATIP, were weakly, but not significantly, inhibitory to neurite growth. Our analyses saw stronger effects (shorter neurites and more condensed chromatin) with the overexpression of LIMK2 and LRRN1 (see also Supplementary Table 3). Finally, Simpson et al. (2008) performed an RNAi screen of kinase and phosphatases genes, assaying the ability of a monolayer of cells from a breast cancer cell line to heal after wounding. They found siRNAs targeting several genes that either potentiated or inhibited the migratory ability of the cells.

Comparing the results of these siRNA screens with each other, it is interesting to note that not a single gene appeared in all three. In fact, only three genes were hits in two of the three screens: BCAR3, LIMK1, and the JNK kinase MAP2K7. The lack of overlap in these RNAi screens supports the notion that such large-scale knockdown experiments require special analysis of the raw data for proper interpretation (Sacher et al., 2008). In our own unpublished experiments using siRNA and shRNA approaches, we have found it much more difficult to knock down protein expression in primary neurons than in cell lines. This is consistent with reports using transgenic mice expressing shRNAs; knock down in the CNS is less effective than in other tissues (Sasaguri et al., 2009). The slow and variable loss of protein expression using RNAi in neurons makes it a problematic strategy in a screening context. A recent high-content analysis of a genome-wide RNAi screen found that the majority of siRNAs have off-target effects (Collinet et al., 2010). Consequently, using RNAi in primary neurons as a screening approach will require much better validated libraries and assays that use novel approaches to allow protein expression to be knocked down before neurite growth starts (Davare et al., 2009). It is therefore critical to use alternative methods, such as overexpression.

Conclusion

Our studies have identified a large number of kinases and phosphatases, as well as structurally and functionally defined families of these proteins, that affect neuronal process formation in specific ways. We have provided an analytical
methodology and new tools to analyze functional data, and have implicated genes with novel functions in neuronal development. Our studies are an important step toward the goal of a molecular description of the intrinsic control of axodendritic growth. Downstream validation studies will be important to perturb the function of other members of the signaling cascades we have implicated, and to examine the functions of these genes in other neuronal types.

Materials and methods

Plasmid cDNA library

A glycerol stock cDNA library from the NIH MGC (Gerhard et al, 2004) in 96-well format was purchased from Open Biosystems (Thermo-Fisher, Huntingville, AL) that included IRAT (human) and IRAV (mouse) clones. The library was replicated and the original and daughter plates stored in −80° degree freezers, sealed with foil tape (Corning Costar 6570, Lowell, MA). The Gene Ontology (http://amigo.geneontology.org/) was consulted to determine an up-to-date list of kinases and phosphatases. Gene ontologies for ‘Kinase Activity,’ and ‘Phosphoric Ester Hydrolyase activity’ were used to generate a list of kinases and phosphatases. During the cherry-picking process, a set of glycerol stock plates into the kinase-phosphatase sublibrary, which Qiagen BioRobot-3000 (Germantown, MD) picked clones from the glycerol stock plates into the kinase-phosphatase sublibrary, which totaled nine 96-well plates. During the cherry-picking process, a set of 10–12 plates were thawed, wiped, uncapped, and the foil cover carefully removed. Disposible tips were used to inoculate media in deep-well 96-well blocks. During the process, glycerol stocks for mCherry (Shu et al, 2006) were also picked and inoculated into specified control wells on the plate. Other wells were left empty, to facilitate non-transfected controls and to allow other clones to be tested.

Plasmid preparation

QIAprep 96 Turbo BioRobot Kit (Qiagen 962141) was used to produce transfection quality plasmid. Briefly, two deep-well blocks from the kit were filled with 1.4 ml Terrific Broth (Invitrogen 2271), Carlsbad, CA) with 150 μg/ml ampicillin (Invitrogen 11539) in each well. A 96-pin replicator tool (Nalge Nunc 250520, Rochester, NY) was used to inoculate the thawed glycerol stock plate into fresh media. Plates were incubated for 20–24 h at 37°C. The plates were spun down serially such that the pellets were overlaid and concentrated. The pellets were resuspended in Qiagen’s ‘P1’ buffer, and the manufacturer’s instructions were followed. Elution was performed at room temperature, with 120 μl of endotoxin-free water. Plasmid concentration was analyzed with NanoDrop spectrophotometer (Thermo Fisher, Wilmington, DE). Plasmid was generally purified at 300 ± 10 ng/μl with an average 260/280 ratio of 2.8. If plasmid concentrations were below 300, plasmids were concentrated by isopropanol precipitation.

Kinase-dead PRKACA mutant

The kinase-dead mutant of human PRKACA was obtained by substituting lysine at position 73 for alanine (G→A substitution in AAG triplet coding for lysine). This was achieved by PCR using the QuickChangell site-directed Mutagenesis Kit (Strategene) with the following primers: K73A sense 5′-GGAACACATATGGCATCGAGATCCT CGACACACA-3′; K73A antisense 5′-TGTTGCGACCACGTCGACAGCCA TAGTTGTTCC-3′. The resulting construct was verified by restriction analysis using the newly created Ncol site (underlined in the sense primer) and sequencing.

Neuronal cell culture

Embryonic hippocampal culture has been described earlier (Goslin et al, 1998; Oliva et al, 2006). Briefly, adult pregnant Sprague-Daley rats were euthanized by CO2 and the E18 embryos were dissected in fresh Hibernate medium (BrainBits, HE-Ca 500, Springfield, IL) supplemented with B27 (Invitrogen 17504). Isolated hippocampi were transferred to Hibernate media without B27 and incubated for 15 min at 37°C with 0.25% Tryspin (Invitrogen 25300), in the presence of DNaseI at final concentration of 0.5 μg/ml (Sigma D5052). The tissue was then washed five times with the same medium supplemented with B27 and triturated until no clumps were visible (about 5–10 times). Dissociated neurons were counted and used for transfection during the next 2 h.

Transfected and non-transfected neurons were grown in 96-well plates (Perkin Elmer, 6005182, Waltham, MA) coated with 10 μg/ml Poly-D lysine (Sigma P7886), and 10 μg/ml laminin (Sigma L2020). When cells were plated on PLL (Sigma P2636) alone as a substrate, the concentration was 1 mg/ml. Enriched neurobasal medium (ENB), modified from Meyer-Franke et al (1995) included Neurobasal (Invitrogen 12348), penicillin/streptomycin, insulin (Sigma 16634 5 mg/ml), sodium pyruvate (1 mM), transferrin (Sigma T1147 100 mg/ml), BSA (Sigma A4161 100 mg/ml), progesterone (Sigma +P8783 60 ng/ml), putrescine (Sigma P7505 16 mg/ml), sodium selenite (Sigma S5261 40 ng/ml), triiodo-thyronine (Sigma T6397, 1 ×), L-glutamine (1 mM), N-acetyl cysteine (Sigma A8199NAC, 5 mg/ml), and B27. Media was also supplemented with CNTF (10 ng/ml).

Transfection

Transfection of embryonic hippocampal neurons was accomplished using the Amaza 96-well ‘Shuttle’ nucleoporation system (Lonza, Walkersville, MD) following the manufacturer’s instructions. Briefly, the Amaza 96-well nucleoporation plate was loaded with the mixture of 75,000 neurons in 20 μl of Amaza transfection solution, and 400 ng of total DNA (including mCherry reporter with ratio plasmid to reporter 6:1) in a volume of 2 μl. The rat neuron transfection, ‘high-efficiency’ program was used, and the neurons were recovered with 80 μl of ENB + HEPES (20 mM, Invitrogen 15630). Several control plasmids were used including pSport mCherry (reporter alone) and pSport CAT, a plasmid with no CMV promoter (chloramphenicol acetyltransferase gene). Cells were then plated at two different densities—8000 and 12,000 cells per well. Transfection efficiency was validated by cotransfecting mCherry with plasmids from the library containing myelin basic protein, vimentin, GFAP, and NCAM cDNAs and subsequent identification by antibody staining (data not shown).

Fix/stain

Neurons were incubated in 37°C 5% CO2 incubators for 48 h. Plates were removed and immediately fixed with room temperature 4% PFA, 4× PBS, in PBS by removing 50 μl of medium and underlaying 120 μl of fixative for 30 min. Plates were rinsed with PBS and stained with anti-IIIIb-tubulin (Aves Labs, TUJ) and Hoechst dye (Invitrogen 33432).

Imaging/tracing

Cellomics KineticScan Reader (Thermo Scientific Cellomics, Pittsburgh, PA) was used to automatically image nine fields in each well of the plates at ×10 magnification in three different channels for nuclear staining (Hoechst), neurite staining (βIII-tubulin), and the reporter gene (mCherry). Images were traced automatically using the Neuronal Profiling Bioapplication version 2.x. For most of the downstream analysis, only the Transfected + and Neurite + populations were analyzed. Reported neurite measures are listed in Supplementary Tables 1–11. Other parameters of high variability in the controls. Supplementary tables include complete results. Images and other data sets are available on request.

Overview

Neurons were transfected and plated in six replicate wells for each cDNA plasmid, at two different densities. Two independent experi-
mental replicates were performed for each of the clones. The clones that changed the morphology of the neurites reproducibly were transfected a third time. Over 10,000 wells were imaged and analyzed, each with nine fields, and one field from each well was manually annotated to check the quality of the automated tracing. The annotation identified acute errors that led to the images being removed from the data set (mostly because of poor focus). The resulting data were normalized (see below) so that the resulting value indicated difference from control for each parameter.

Analysis

Data aggregation/storage

Raw data were managed by the Cellomics Store, which consists of an SQL database and a network-attached fileserver (HP). Raw data consisted of metadata associated with scanning and tracing (exposures, focus offsets, thresholds), raw images and the results of the tracing. In addition, cell and well level data were exported and stored on a separate fileserver, organized by experiment with accompanying Excel tables listing how particular wells were treated.

Spotfire DecisionSite (Tibco) was used to associate the treatment variables (which plasmids were transfected in which wells) and perform basic quality control, including checks for tracing errors, low-and high-density wells, cell clumps, and plating errors.

Tracing quality control

Tracing was validated several times by comparison with NeuroLucida tracing (MicroBrightField, Williston, VT). For validation of the tracing in the screen, an image of the tubulin channel from the first field of every well in each plate (over 10,000 images) was exported as a jpeg. Then, three annotators ran a custom C# program on their desktop computers that displayed the image and gave several options as to the quality of the focus, threshold, tracing, and cell density. The combined data from the annotators were used to either retrace plates or eliminate wells from the analysis.

Transfection threshold

By examining the red fluorescent intensity of the neurons that were not transfected with plasmid, a background amount of intensity was established using Spotfire DecisionSite, and used as a threshold for the classification of neurons as transfected. Within each experiment, a transfection threshold was defined using the distribution of the average red intensity (AI) of each cell in control neurons that were not subjected to nucleofection. This distribution was near to log-normal. Treated cells with a log(AI) higher than mean + 2 s.d. of the control log(AI) were considered transfected. Only transfected cells were included in the analysis.

Normalization

The morphometric data of transfected cells in each treatment were normalized with respect to a control within the same experiment. We have observed that experiments, defined as the sequence ‘isolation of neurons/transfection/culture/staining/imaging,’ are a major source of variation. The controls are neurons that were subjected to transfection with the transfection marker and the plasmid pSPORT CAT as well as the reporter mCherry. After normalization, the variables for a given experiment i were aggregated across the N experiments, disregarding its statistical significance given that it is not dependent on sample size and dispersion.

To estimate the statistical significance of the effects produced by any treatment, a different normalization was used. It is a t-value that is aggregated as a simple sum across the N replicated experiments. The statistical significance of this sum is computed from the convolution of N Student distributions each with ntransf + ncontrol – 2 degrees of freedom.

Linear model

A generalized linear model for the NTL as a function of experiment and treatment was built to evaluate the quality of cotransfection-based cell selection, the normalization scheme and the inter-experiment variability.

\[ \text{NTL}_{ij} = C + E_i + T_j + ET_{ij} + e_{ij} \]

where C is the overall average NTL, \( E_i \) represents the effect of the ith experiment, \( T_j \) represents the effect of the jth treatment, and \( ET_{ij} \) represents the interaction between \( E_i \) and \( T_j \). This model was fitted to the NTL data from four (\( i = 1 \ldots 4 \)) controls, pSPORT CAT, RFP, KLF4, and KLF7 across eight experiments (\( j = 1 \ldots 8 \)). KLF4 and KLF7 are genes that systematically affect neurite growth (Moore et al., 2009). The variable NTL was analyzed in its original form and after the two normalization procedures for both transfected and non-transfected cells analyzed independently. Thus, the model was fit to six different data sets, each of them with 64 data points (8 × 4 × 2 densities), producing 32 degrees of freedom for the error \( e_{ij} \) that is assumed to be independent and identically distributed with mean=0.

Correlation analysis

A Pearson correlation was applied to normalized data. Bootstrap resampling was used to construct a CI for the correlations using custom software written in C# with Microsoft Visual Studio 2008. During bootstrapping, pairs of values were taken by replacement n times to reconstruct a bootstrap replicate of the entire population. That process was then repeated 10,000 times, and the correlation coefficient was stored for each run. CI is reported as the top and bottom 2.5% correlation coefficients.

Neuron survival

Nuclear size and intensity of DNA staining (Hoechst) have long been used as indicators of the live/dead status of cells. Small and bright nuclei are the result of heterochromatin condensation, a hallmark of apoptotic cells death. Big and dim nuclei correspond to cells that are alive. We use this distinction to measure the cell survival of our cultures transfected with a variety of cDNAs, placing emphasis on those cDNAs that we found to significantly reduce neurite outgrowth.

In most instances, the clusters of live and dead cells in the nuclear area versus intensity were easily distinguished. Semi-supervised classification was implemented using three categories (live, dead, out of range) using a custom script in MatLab and was run on data on a plate-by-plate basis. The survival rate varied as a function of the transfection and neurite-bearing status. The RFP + cells with neurites (\( N + RFP + \)) showed the highest survival rate.

As a measure of whether survival rates in treatments reducing neurite growth were significantly lower, cells in which cDNA for RFP was transfected with 240 ng of DNA were used as controls (same amount of cDNA used in the screen). Z factors were computed within each experiment using the mean and the s.d. of the control. The results within each treatment per replicate experiments were averaged and significance was computed using a permutation test. Supplementary
Figure 3B shows the signed significance for the effect of genes on survival. The signed significance is the absolute value of the log10 (P-value) multiplied by the sign of the Z factor. A signed significance of $\leq -2$ means that survival of the treatment is significantly lower than the control.

Bioinformatics

Phenotype clustering

After averaging values from genes that were represented by two species or multiple clones, phenotypic classes were determined by taking a discrete measure of significance ($-1$ significantly below control, 0 not significant, and 1 significantly above control at $P \leq 0.05$) and clustering to form distinct groups based on neurite average length and primary neurite count. Clustering was performed on the average values for neurite average length, primary neurite count, and branching with the main groups ‘stratified’ by the prior discrete clustering. The hierarchical clustering algorithm used normalized phenotypic data as well as the statistical status of each gene, but did not consider which genes were transfected or to what classes they belonged. This led to the arrangement in Figure 3. Hierarchical clustering was performed in Spotfire DecisionSite using clustering algorithm UPGMA and Euclidian distance.

Phylogenetic trees

Genes were grouped into five main categories, based primarily on their gene ontology information, but additionally hand curated to ensure higher accuracy. Most protein kinases were considered only if they were listed in one of the kinase references (Manning et al., 2002; Caenepeel et al., 2004). A few genes had ambiguous classification as they interacted with both kinases and phosphatasas or had reported activity but no obvious enzymatic domain, etc. The process of tree creation is outlined graphically in Supplementary Figure 5. Protein sequence was obtained from the gene’s Entrez Gene (http://www.ncbi.nlm.nih.gov/sites/entrez?db=gene) entry, and loaded into EBI’s ClustalW2 (http://www.ebi.ac.uk/Tools/clustalw2/) (Thompson et al., 1994; Larkin et al., 2007) for each group separately in FASTA format. The output of the cluster algorithm is a dendrogram file, which represents the hierarchy in a nested text format [i.e. (Parent(Child A:0.2,ChildB(GrandchildA:0.13,GrandchildB:0.1:0.3))]]. The dendrogram file is read into custom software, which parses the dendrogram and represents it as a binary tree, which can be drawn automatically in CorelDraw (X3, Corel Corporation, Mountain View, CA) as a radial, centered tree. Branch lengths in the tree correspond to the log of the distance as reported by ClustalW2. The angles of the branches and the order of the two segments from a branch are arbitrary and cosmetic. Some branches have multiple markers, indicating different clones that were tested for that gene (human and mouse as well as clone variants of the same species). Other branches have no marker, indicating one of the following: the gene was screened but the replicates failed, there were too few cells to analyze (possibly a viability defect), or failed tracing or image validation.

Phylogenetic gene cluster analysis

Phylogenetic relationships (from above) were transformed into a table where the columns represented nodes of the tree (except the leaves), and records were the genes. Values were binary representations of a gene’s presence in that node (the column representing the node would get a 1 for presence or 0 for absence). That table was imported into Spotfire and hierarchical clustering (using correlation and UPGMA) was performed. A custom C# program was run that sliced through the hierarchy at any number of levels (termed tiers). The program operates in one unit at a time—a ‘node set.’ Each node set would have some number of genes and the total number of genes would be represented by the total node sets across the tier. For each node set, the values for a particular morphological parameter from the member genes were averaged. The data were represented graphically as a cluster heat map, where each tier is a row in the graph, and a node set is a rectangle, colored by its average value. Statistics fit the cluster averages with a bootstrap sample from the entire data set. Significances are indicated by diamonds and asterisks. Diamonds indicate ‘spiked bootstrap’—the family had a significant average when considering it against random picks always including the maximum or minimum value of the parameter (similar to first inverse jackknife) (Efron and Tibshirani, 1994). Asterisks indicate the family had a significant average, even after correcting for multiple comparisons per tier by Benjamini/ Hochberg methods (Benjamini and Hochberg, 1995) ($z = 0.05$). The analyses were run separately for each parameter tested, and significant values were summarized.

Pathway gene cluster analysis

Pathway information was garnered from Entrez Gene through linking to Reactome (http://www.reactome.org/) and KEGG, (http://www-genome.jp/kegg/pathway.html) (April 2009). Pathway annotations were found for 49% of the genes. KEGG not only provides pathway annotation, but also has a hierarchical representation of the pathways, so was chosen for further analyses. The hierarchical categories, subcategories, and pathways themselves were columns of a table, and genes were in each row. If a gene (e.g. IKBKB) was a member of the ‘Pancreatic cancer’ pathway, then it was assigned a 1 in that column, a 1 in the column ‘5.1 Cancers,’ and a 1 in the column ‘5 human diseases.’ In this cluster analysis, genes were allowed to be duplicated, so IKBKB had other rows in the table, and was assigned with the information for other pathways as well (chronic myeloid leukemia, prostate cancer, small cell lung cancer, type II diabetes mellitus, apoptosis, B-cell receptor signaling, T-cell receptor signaling, Toll-like receptor signaling, adipocytokine signaling, insulin signaling, MAPK signaling). Pathways with fewer than two genes were excluded. Average values were taken for each group, and displayed on a heat map as before. Here, a small square symbol indicates the family had a significant average but failed to be significant after corrections, and diamonds indicate significance compared with bootstrap replicates always including the maximum or minimum value of the parameter being tested (spiked bootstrap). When moving to lower tiers, genes were not allowed to be duplicated within a group, but were allowed to be duplicated between groups.

Supplementary information

Supplementary information is available at the Molecular Systems Biology website (http://www.nature.com/msb).

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Conflict of interest

The authors declare that they have no conflict of interest.

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