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METHODODOLOGY

A Novel Multi-Scale Modeling Approach to Infer Whole Genome Divergence

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Abstract: We propose a novel and simple approach to elucidate genomic patterns of divergence using principal component analysis (PCA). We applied this methodology to the metric space generated by M. musculus genome-wide SNPs. Distance profiles were computed between M. musculus and its closely related species, M. spretus, which was used as external reference. While the speciation dynamics were apparent in the first principal component, the within M. musculus differentiation dimensions gave rise to three minor components. We were unable to obtain a clear divergence signature discriminating laboratory strains, suggesting a stronger effect of genetic drift. These results were at odds with wild strains which exhibit defined deterministic signals of divergence. Finally, we were able to rank novel and previously known genes according to their likelihood to be under selective pressure. In conclusion, we posit PCA as a robust methodology to unravel diverging DNA regions without any a priori forcing.

Keywords: principal component analysis, adaptive evolution, genetic drift, multi-scale modeling, speciation, reproductive isolation

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Introduction
One of the most challenging tasks in molecular evolution is to identify genetic loci that may contribute to phenotypic and genotypic divergences. In general, genetic divergence can be classified into deterministic and stochastic parts. The deterministic part of divergence may convey the beneficial effects forward to the next generation, whereas the stochastic part of divergence may represent neutral genetic drift. For example, selective sweep describes a process in which neutral sites physically linked to an adaptive mutation can result with a skew in the distribution of the allele frequency. In general, it is possible to consider such spectra of genetic variation as being deterministic because they can be traced back to a selective process. Tests for neutrality can be inferred by examination of genetic differentiation over short periods of time (ie, the \( F \) statistics). Loci that are found to exceed an empirical \( F \) threshold are considered candidates for stabilizing or directional selection. In contrast, there are other methods that examine divergence after a longer period of evolution and mostly relies on the relationship between the synonymous and nonsynonymous mutations (eg, the \( d_{s}/d_{a} \) substitution rates or the McDonald–Kreitman test). In principle, those methods assume that the same rate of change can be considered as neutral, while significant deviation from neutral evolution can be inferred as negative or positive selection. It is interesting to consider that all the above methods are built upon a single-gene (locus) basis without taking into explicit consideration the inherent multidimensional character of selection.

The house mouse \( M. \) musculus and its closely related species \( M. \) spretus are perfect model organisms for studying evolution. \( M. \) musculus includes three main groups of isolated wild subspecies (\( M. \) m. domesticus, \( M. \) m. musculus and \( M. \) m. castaneus), which diverged from the last common ancestor around 1 million years ago and migrated into three distinct continents following allopatric separation. The potential of using the house mouse for evolutionary studies is also attributed to the presence of extant laboratory strains. These strains are very well characterized from the phenotypic point of view, and, due to strictly controlled laboratory conditions, natural selective processes are virtually absent. The genomic background of these strains is thought to be composed of “mosaic” fragments of high and low differentiation (inter and intra sub-specific origin with correspondence) following historical domestication events of the \( M. \) musculus subspecies. Even if their pedigree is unknown, one may postulate that since the genetic makeup of mosaic strains was not shaped by past natural events but due to random human artificial breeding, the signal derived from adaptive evolution should decrease relative to the wild strain, whereas the effect of drift should increase. This is also due to random-sampling matings (or “founder effect”) of individual mice from natural breeding as well as their relative small population size.

Principal component analysis (PCA) is a multidimensional statistical technique particularly suited for optimizing the amount of biologically relevant information that may derive from genome scans of DNA polymorphism. PCA has already demonstrated its effectiveness in singling out features that are invisible to single-gene oriented techniques in gene expression analysis. In addition, PCA is a common approach in the field of population genomics and is constantly used to unravel recent demographic radiation of populations. When applied to a distance space, PCA is able to project the system into a component space in which the main “directions” of variance are easily interpreted. This is especially evident when distinct order parameters are present, directing the distances of between elements to size and shape components. While the size (first principal component, all positive loadings) captures the across genome correlated distances to the reference, common to all sampled objects, shape (minor components, both positive and negative loadings correspondent to a balance among different order parameters) represents the modulation (size independent) of the distance space among the sampled objects.

There are several examples of how deterministic signal can be differentiated from stochastic noise by deconvolution of the two mutually independent modes using PCA. These include fields of plate tectonics, meteorology and oceanography, or global positioning (GPS). In this study, we applied PCA methodology to distinguish between deterministic and stochastic drift patterns of genomic divergences using distance spaces derived from around eight million distinct patches of DNA. We found that the distantly related strain \( M. \) spretus could be used as an optimal
reference for the computation of genetic distances for the entire *M. musculus* genealogy. Moreover, by using the score space of the major principal components we were able to distinguish between minor fractions of deterministic signals embedded within the stochastic noise (genetic drift or metric errors).

Our results illustrate that while the first principal component (PC1) captured the speciation signal between *M. musculus* and *M. spretus* with 91% of the variance, the orthogonality between the components explained the differentiation between wild and lab strains with 3% of the variance. It will be demonstrated that wild strains that can be differentiated by PCA possess abundant fractions of deterministic signals (loci) that underscore significant association with functional groups. On the other hand, no such localized regions of divergence could be detected in the lab strains, supporting the proof of concept that selection of deterministic signals may play some significant role in evolution. In sum, the results to be described, indicate PCA as a potentially useful methodology for “deep sequencing” based evolution studies. There is thus a rational basis to explore the characteristic nature of evolutionary dynamics while minimizing the initial assumptions placed on the data.

**Methods**

**Dataset**

Mouse SNP data were obtained from the Sanger web site. We used a SNP list from 13 mouse laboratory strains (129P2, 129S1/SvImJ, 129S5, A/J, AKR/J, BALB/cJ, C3H/HeJ, C57BL/6NJ, CBA/J, DBA/2J, LP/J, NOD/ShiLtJ, NZO/HiLtJ), 3 wild-derived *M. musculus* subspecies (CAST/EiJ [*M. m. castaneus*], WSB/EiJ [*M. m. domesticus*], PWD/PhJ [*M. m. musculus*]) and one wild-derived *M. spretus* species (Spretus/EiJ). All data were stored and annotated using a MySQL database with custom Perl and Java programs. A full dataset of exons and introns was obtained using the BioMart tool from Ensembl, version 61 (Ensembl, Cambridge, UK).

**Data analysis**

The Sanger Institute has recently established a new mouse resequencing project in order to investigate the phylogenetic origin of laboratory mouse strains and to make a comprehensive investigation of the mouse genetic variation using a list of approximately 65 million high confidence SNP calls. This is the dataset we used as a case for evolutionary study and to validate the reliability of our PCA methodology. We first partitioned the genome into equal lengths of genetic loci by using a sliding-window technique. In order to obtain the best resolution for the minimum reliable window size, we chose various windows lengths of $L \in \{50, 200, 500, 1000, 2000\}$ for all autosomes and the X chromosome. For each window length, $l$ (or genetic loci), we calculated the average number of pairwise SNP differences ($\pi^*$) between laboratory and *M. musculus* subspecies and *M. spretus*. We then set the data matrix, $G_j$, of $j = 17$ columns representing the different strains in terms of distance from *M. spretus* computed for the different windows as shown in formula 1.

\[
G_{i,j} = \begin{bmatrix}
\pi_{1,1} & \pi_{1,2} & \cdots & \pi_{1,j} \\
\pi_{2,1} & \pi_{2,2} & \cdots & \pi_{2,j} \\
\vdots & \vdots & \ddots & \vdots \\
\pi_{l,1} & \pi_{l,2} & \cdots & \pi_{l,j}
\end{bmatrix}
\]

Formula 1: Distance matrix.

Each data matrix, $G_{l,p}$, was correspondent to $l \in \{18, 8, 4, 2, 1\}$ million statistical units (rows) of distinct measurements of the genetic distances in consecutive order along the chromosomes. For example, the data matrix of window size $l = 200$ contained $i = 8$ million genetic loci, each one represents the genetic distances between all 17 *M. musculus* with *M. spretus*. It is important to note that since *M. spretus* was the reference strain, the distance matrix, $G_{jp}$, does not contain *M. spretus* as a variable.

Our first intention was to determine the smallest locus size that could still show a minor decay of the between strains correlation despite the obvious discretization of the data due to small sampling of mutations. To do this, we calculated the pairwise correlation coefficients between the wild strains for each distance matrix, $G_{l,p}$, separately for each window size and identified that the between strains correlation had a major decay when the median locus size was shifted between $l = 200$ and $l = 50$ while the correlation coefficient was stationary when $l > 200$ (Fig. 1).

As expected, the X chromosome consistently showed a lower divergence than the autosomes for all strains.
We observe that the distance between *M. musculus* and *M. spretus* is represented by size and shape components. The size component (major component PC1, all positive loadings) captures the across genome correlated distances among the *M. musculus* group and *M. spretus*, common to all *M. musculus* strains. The shape component (minor components PC2, PC3, PC4, - positive and negative loadings) represents the modulation (size independent) of *M. musculus*—*M. spretus* distances within the *M. musculus* group. The possibility of reliably discriminating between loadings (strains) provides essential knowledge of the genetic distance space represented by the eigenvector of each principal component. Thus, it becomes feasible to further classify each genetic locus into signals of high or moderated divergence without the need to rely on any a priori information.

**Identifying deterministic signals of divergence**

Having demonstrated the efficacy for strain discrimination of the loading space, we next concentrated on the component score space to extract regions endowed with the highest divergence signals. First, PCA was calculated for each chromosome independently, and the score profile of each principal component was converted into a discrete binary variable \( b = \{0, 1\} \) where “1” substituted for any absolute score of a locus greater than \( 2\sigma \). Since scores represent the magnitudes of the relationship between each datum of the manifest and latent variables, these genetic loci were considered as extreme score loci (ESL) and hotspot candidates for divergence. In contrast, “0” was assigned to the remainder of the dataset as regions of less importance. Second, each ESL was classified as intronic, exonic, or intragenic regions for further analysis. Since PCA is an unsupervised method, each ESL could be considered as a reliable classifier for divergence hotspots. In order to further assess and distinguish between genetic loci of deterministic or drift signals, the sliding windows technique for 30 consecutive loci (~6000 bp) was applied. In each step, significant clusters of ESL (cESL) were checked for by using hypergeometric distributions. Significant regions were observed when \( P_R < P_B \), where \( P_R \) denotes the \( P \) value of a tested region and \( P_B \) denotes the Bonferroni threshold (see Formula 2).
Formula 2: Probability mass function of hypergeometric distribution.

\[ P(X = x) = \binom{k}{x} \binom{M - K}{N - x} \binom{M}{N} \]

- \( x \), the number of ESL in each one of the tested windows.
- \( M \), the total number of windows in each chromosome.
- \( K \), the number of ESL in the entire chromosome.
- \( N \), the number of windows for each calculation (constant 30 genetic loci).

From each clustered region (cESL), the corresponding ESL was extracted for further analysis whereas the rest of the genome was disregarded (ie, ESLs that were not clustered together). We assumed that those regions were hotspot candidates for evolution or regions with linkage to adaptive regions.\(^7\) We also assumed that dispersed ESLs (ESL that were not clustered together) may represent regions of past episodic selection or drift. In order to test whether our native dataset contained evidence for localized regions of divergence (therefore proving strong linkage between neighbor loci), we permuted the original dataset for each chromosome so the hierarchical order between neighbor loci could not derive from their physical linkages. We then reapplied PCA on the permuted dataset relying on the robustness of PCA to keep the cumulative distribution of the scores along the eigenvector unchanged, but with different order along the chromosome. We also checked that the permutation of the native data matrix had no effect on the structure of the loadings space. Plots were then made of the permuted dataset and the native dataset with the Bonferroni threshold. It was found that the native dataset contained ESL with tide linkage compared with linkage-free relationships between loci in the simulated dataset (Fig. 2). Interestingly, while PC1 to PC3 contain more localized clusters of cESL, PC4 clusters tended to be more overdispersed across the genome.

The ability to partition the entire genome into small genetic loci (200 bp) together with the observation that some of the genetic loci are particularly more reliable and localized within specific regions allows us to classify significant genetic loci into exonic, intronic, and intergenic locations for each principal component with much better accuracy. Thus, by employing PCA strategies it is possible to gain a better association between the loading space, the distance space, and their potential functions.

**Results**

**Size component (PC1) loading distribution interpretation**

The possibility to reduce the dimensionality of the system genetic variation into orthogonal principal components allowed us to explore the flux of genetic variance that explains different trajectories of genomic evolution. All genetic distances in our model were computed by referencing various \( M. \) *musculus* strains with \( M. \) *spretus*. Since the between species distances are much bigger than those of the within species, speciation and reproductive isolation may be inferred from the presence of distinct size (between species) and shape (within species) components (see Method section).

As expected, PC1 captured by far the majority of the correlated distance between the two species (92% of the explained variance) with constant loadings scores for all \( M. \) *musculus* group (wild-derived and laboratory strains, loadings \( \approx 0.3 \), Fig. 3). This implies that the major portion of the genome-wide divergence profile between the \( M. \) *musculus M. spretus* groups is extremely invariant. This reflects the genetic distance flux corresponding to the speciation between \( M. \) *musculus* and \( M. \) *spretus*. However, despite the fact that most of the profile of genetic distances between the two species is highly correlated, the within species distance is not coincident and can vary between genetic loci of the different mouse strains. The capturing of 92% variance in PC1 between the two species along their entire genomes confirms that \( M. \) *spretus* is an outlier species with respect to \( M. \) *musculus* strains. However, PC1 appears substantially invariant among different strains (Fig. 3), implying that the within species variation has a different nature (not only a different entity) with respect to the between species one.

**Minor components loading interpretation**

In contrast to the invariant profile of genetic diversity carried by PC1, all minor components combined (PC2-PC4) explained only 8% of the genetic diversity, but
still faithfully reflected the modulation of variability within the *M. musculus* group. Regarding genetic variability, we examined only the first three minor components (PC2, PC3, and PC4), which explained about 3% of the variability, which makes a clear discrimination between the wild and the laboratory strains. The other components were disregarded as noise.

Our first observation from the loading space of the minor components revealed clear differences between wild-derived and laboratory strains, which are consistent with their varied ancestral origins and past lifestyles. PC2 (2.5% variance) explains the divergence between *M. m. musculus* and *M. m. castaneus* (loading = 0.6) from *M. m. domesticus* and the other laboratory strains (loading = −0.1, Fig. 3). Such a clade of wild and laboratory strains that are clustered together in the same principal component illustrates that the laboratory mouse strain shares a larger genetic profile with *M. m. domesticus* than with *M. m. musculus* and *M. m. castaneus*. This finding is consistent with past studies identifying *M. m. domesticus* as the most dominant founder for the laboratory strains\(^{17,18,32}\) and is a proof of concept that the loading space correctly mirrors phylogenetic relationships among *M. musculus* strains.

PC3 (1.1% of variance) highlighted an evolutionary pattern similar to PC2. Whereas PC2 gave the same
weight for the genetic variance shared by laboratory strains and *M. m. domesticus*, PC3 discriminated only wild-derived strains (*M. m. castaneus*, *M. m. musculus*, loading = 0.6 and −0.6 respectively). *M. m. domesticus* and the laboratory strains had minimal or null effects (loadings ≈ 0). Then, in contrast to PC2 and PC3, PC4 (1% variance) gave rise to a clear discrimination between two groups of laboratory strains 129 (129P2, 129S1/SvImJ, 129S5) and AKR/J versus the rest (loadings ≈ 0.4, loadings ≈ −0.2). In these cases, the three wild-derived species were kept horizontal to the axes of the principal component space (loadings ≈ 0). These specific results evoke different explanatory hypotheses: (1) PC4 gives rise to a unique adaptive process that is captured neither by PC2 nor PC3, meaning that it has a low likelihood of stemming from the large signal reported from wild-derived discrimination in PC1-PC3; (2) adaptive signals are possibly derived from an unsampled ancestor (proof is beyond the scope of this study); and (3) signal arises from genetic drift (validation requires future downstream analyses).

Having demonstrated the biological relevance of both the loading spaces and extracted components, we are allowed to explore the score space, that is, the values assigned to different genetic loci by the above described components. As an aside, it is worth noting that PCA is a fully unsupervised approach and the strain discrimination emerged a posteriori from the analyses.

Underdispersion of ESL in PC4

We calculated the dispersion index of ESL for each one of the four component scores as indications of their randomness of distribution for each cluster (Fig. 4). In general, both the first component (PC1) and the minor components (PC2-PC4) included a similar number of ESLs attributed to their distribution profiles with common dispersion indexes approximating Poisson distributions $D \approx 0.97$. Even though cESL tended to cluster within specific hotspot regions of evolutionary importance, it is essential to understand the level of abundance of cESLs for the

![Figure 3. Loading space of PC1 to PC4.](image)

**Notes:** The figure illustrates constant loading profile for all *M. musculus* strains for PC1 and hierarchical modulation of the minor components between the *M. musculus* strains. It is apparent that while PC2 and PC3 discriminate wild derived strains, PC4 discriminates two group of laboratory strains which could be segregation driven by genetic drift.

![Figure 4. Cumulative distribution of dispersion index units of ESL across window groups of 30 genetic loci for the first four principal components.](image)

**Note:** PC4 tends to group fewer ESL within regions than observed in PC1- to PC3, thereby affirming that those regions are under recent divergence.
different clusters. With this aim, we used sliding window approach \( (w = 30 \text{ genetic loci}) \) and measured the dispersion index of cESL within each window. We then summarized the cumulative distribution of dispersion scores for all windows as a benchmark for the level of dispersion of cESL within each window. Interestingly, we found that clusters within the first three principal components are less enriched with ESL and tend to exhibit higher dispersion values. In contrast PC4 shows a significant downshift of dispersion along sequential windows. Thus, despite the fact that PC4 clusters tend to be more dispersed along the genome (Fig. 2), those regions are more enriched with cESL. In this case, PC4 shows more dispersed regions of diversity with more regions of recent divergence compared with the other minor components (PC2 and PC3). This result is in line with the loading profile of PC4, which makes a clear discrimination between the different laboratory strains tested. Therefore, we propose PC4 as a candidate component that captures the flux of variation correspondent with genetic drift whereas PC1 to PC3 may be more affected by the selective process.

Genetic divergence analysis

While the loading space highlights latent variables explaining whole genomic evolutionary forces, the score space presents a more precise view of genomic regions involved in specific evolutionary processes. Therefore, prior knowledge of the loading profile for each principal component can be supported by detailed analyses of genetic loci as possible candidates for selection or drift. We have used the list of cESLs from each principal component (see Methods) and classified each genetic locus into exonic, intronic, and intragenic regions for further analysis. We focused on cESLs underlining exonic regions from the likelihood they may exert more effects on the function and fitness of the organism. In general, we found that the number of candidate genes increases together with the hierarchical order of the principal components. For example, the number of genes listed in PC4 was significantly higher than the one observed in PC3 and PC2 (Table 1). Those results were consistent with noncoding RNA and pseudogenes. This finding is perfectly consistent with the high dispersion levels of PC4 and opposed to PC1-3 dispersion, which tends to be localized to specific regions. In addition, we found that while exons with extreme PC4 scores were randomly scattered across functional classes, exons with higher PC1-3 scores illustrated a clear association with functional groups.

Table 1. Summary report of cESL across principal components.

| # genes | Exon enrichment | Intron enrichment |
|---------|-----------------|------------------|
|         | # genes | # genes | # genes | # ncRNA | # ncRNA | # pseudogenes | P value | q value | P value | q value |
| PC1     | 100     | 38      | 84      | 8       | 1       | 12            | 1 × 10^{-5} | 1.002   | 0.001   | 0.3     |
| PC2     | 352     | 168     | 293     | 23      | 10      | 44            | 1 × 10^{-6} | 1.002   | 1 × 10^{-5} | 0.02    |
| PC3     | 528     | 332     | 445     | 31      | 22      | 60            | 2 × 10^{-10} | [1]1 × 10^{-7} | 7 × 10^{-9} | [2]1 × 10^{-5} |
| PC4     | 1432    | 970     | 1200    | 110     | 56      | 107           | 3 × 10^{-4} | 0.5     | 2 × 10^{-3} | 0.2     |

Notes: [1]Olfactory receptor; [2]Synaptic transmission; [3]Immune system genes. The number of candidate genes is significantly increased in PC4 compared to the other principal components. As expected, candidate genes within PC4 are scattered randomly across functional groups and strengthens the hypothesis that those genes are likely to be under the effect of genetic drift. In contrast, genes among the first 3 principal components are clustered among functional groups, which underlies a possible mechanism of natural selection.
ing evolution, the fact that intron enrichment occurred only within this class with no evidence for candidate exonic regions points to a neutral divergence with no imposing functional consequences. In addition, the lack of any intron enrichment of olfactory receptors is an artifact of sampling since olfactory receptors are very poor in introns. In contrast to PC2, both intron and exon candidates of PC3 show a clear association with immune system genes. The enrichment is more significant among candidate exons even if those genes are significantly more abundant with introns and suggests that these genes could undergo selection. One of the main characteristics of PCA is that individual principal components are orthogonal to each other. Because of this constraint, one would expect that genetic loci with high contributions to PC2 not be represented very well among the other principal components. However, since the dataset employed in this study focused on a single gene covered by many genetic loci, it is still expected that a certain level of superposition of candidate exon genes among the different principal components would be found. In that case, candidate genes that were found to be shared by different principal components could point to regions that have relevance for the discrimination of different lineages. Alternately, candidate exon genes specific to only one principal component could point to a lineage of specific divergence. In addition, the fact that shared candidate exons genes were represented in different principal components reveals a significant nonrandom selection of genes toward functional classes. In general, most of the candidate genes were unique to corresponding principal components (Fig. 5) with only one gene being common for PC1-3 (Skint5—T cell). We obtained 26 overlapping genes (~15%) between PC2 and PC3, most of which were histocompatible, olfactory, or C-type lectin-containing domains. This group of 26 common genes was significantly more enriched than was expected by chance alone (2 expected shared genes). This is further proof of the accuracy of PCA methodology to locate major genes playing possible roles in divergence. The fact that most of the candidate genes from both components are known to be under strong selective constrains during evolution affirms that PCA is able to decompose the genetic variation into selected regions of divergence without losing the global picture of the latent directions of divergence between various evolutionary scenarios.

**Figure 5.** Venn diagram illustrates common candidate exons genes for PC1 to PC3.

**Notes:** The figure suggests that ~85% from the genes are unique for each one of the principal components. This is expected since the principal components are each orthogonal. However ~15% (26 observed genes) of the exon genes are shared among principal components as extremes. This number is much higher than what expected by chance (2 expected genes) and confirm their essentiality for divergence.

**Discussion**

Recent advances in sequencing technologies allow the acquisition of high resolution sequence information for entire genomes from which knowledge of the dynamics and trajectories of evolution between and within closely related species can be deduced. A better accuracy of the read calls leads to better confidence in SNPs, reducing the likelihood of making false positive and false negative predictions. Since the amount of future data is predicted to grow exponentially, new methods to handle this huge data flow are required. In addition, the fact that most of the existing methodologies are single, gene-centric oriented, they are incapable of taking a higher dimensional, more global approach on data if and when require.

In this study, we have presented a PCA-based methodology to unravel regions of important genetic divergence. We have also demonstrated that an apparent drift component could be identified in a small and artificially-bred population of laboratory animals. This finding was in direct opposition to more deterministic behavior as captured among the principal components of divergence in wild strains. Likewise, we have demonstrated that our PCA approach provided two consistent associated representations of
strains (loading space) and genetic loci (score space). Our results point to a leading common “speciation” first principal component (91% of the explained variance) which implies a very strong correlative structure of the distance between the two species measured at different loci. We found that the score distribution of the first four principal components mirrored the evolutionary trajectories drawn by the loading space following the multi-scale modulation of the genetic distance between \textit{M. musculus} strains and \textit{M. spretus}. It is apparent that while PC1, PC2, and PC3 discriminate wild strains, PC4 discriminates laboratory strains.

By using a clustering approach, we were able to identify hotspot regions for divergence and to characterize each one of the four principal components as potential random drift or deterministic divergence that may indicate selection processes. In contrast to PC4, the first three principal components exhibited more deterministic behaviors, which can be interpreted as: (1) discrimination of wild strains; (2) localized hotspot of divergence (Fig. 2); and (3) significant associations with function (Table 1). We conclude that PC1 to PC3 are explained more by evolution following adaptation, whereas PC4 illustrates evolution by drift with a more neutral effect.

As expected, by using high-resolution analyses of exons from candidate gene loci, enrichment was found in the second and third principal components. Interestingly and unexpectedly, each principal component was enriched with different groups of genes according to their functional groups. While PC2 was associated with the olfactory receptor, PC3 included only enrichment of immune system genes (Table 1). We found especially that the Skint family was remarkably influential by discriminating principal components with a very strong signal from PC3 (which in general was more overdominated with immune system genes, Table 1). Olfactory and immune system genes are known to be under strong selective pressure in the mouse\textsuperscript{13,35,36}. Immune system genes are known to be under strong positive selection in \textit{Drosophila}\textsuperscript{37,38} and, in general, in all vertebrates\textsuperscript{39}. The fact that different gene families exerted major effects on different principal components could be due to the fact that each orthogonal component represents different episodes of weak to high evolutionary pressure.

It will be necessary to validate our evolutionary hypothesis with pre-existing population genetics methods such as the $d_\text{S}/d_\text{S}$ substitution rates\textsuperscript{9–11} or the $F$ statistics. However, this approach is not feasible with our present dataset due to insufficient sequenced animals either to perform the $F$ test or to meet the robust statistical assumptions required by the maximum likelihood framework. However, even though validation with known tools to assess adaptive evolution cannot be performed in the context of our paper, we were nonetheless able to single out a group of genes that are known to be under strong selective constraints. And since these results were obtained without any a priori information, this validates that our PCA methodology be applied to other model systems as a suitable complement to other extant multilocus methodologies.

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\textbf{Author Contributions}

Both authors ER and AG planned the research. ER was mostly responsible for carrying out the analyses. ER did most of the writing, which was edited by an English-language reviewer. Both authors read and approved the final manuscript.

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Figure S1. Correlation of genetic divergence in chromosomes 1 and X. It is notable that authosomes are over-diverged then the X chromosome due to increase amount of meiosis during evolution (A). In addition lab strains illustrate less divergence signals due to more homogenous genetic background (B).

Figure S2. Illustrate is a matrix of Pearson correlation coefficient between studied mouse strains.

Figure S3. Table in excel format represent candidate genes under divergence for the 4 major components (PC1-PC4). (File available from article homepage).