GeneGini: Assessment via the Gini Coefficient of Reference “Housekeeping” Genes and Diverse Human Transporter Expression Profiles

Highlights
- Gini index (0–1) is a convenient means of summarizing inequalities of distribution
- We apply it to two, large transcriptome datasets from tissues and cell lines
- Membrane transporters (SLCs) have unusually heterogeneous distributions
- Low Gini index transcripts make great reference genes; we describe many new ones

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In Brief
The Gini index (coefficient) is used by economists to describe inequalities in wealth distribution in populations and varies between 0 (full equality) and 1 (extreme inequality). We here adopt it to describe, in a simple way, the distributions of expression levels of different genes between tissues or cell lines. We find that uptake (SLC) and efflux (ABC) transporters are more heterogeneously distributed than are members of most other gene families. By contrast, genes with a low Gini coefficient must be stably expressed and can be proposed as reference genes for normalization in expression profiling studies. As judged by this criterion, many previously unidentified reference genes may be proposed.

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GeneGini: Assessment via the Gini Coefficient of Reference “Housekeeping” Genes and Diverse Human Transporter Expression Profiles

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SUMMARY

The expression levels of SLC or ABC membrane transporter transcripts typically differ 100- to 10,000-fold between different tissues. The Gini coefficient characterizes such inequalities and here is used to describe the distribution of the expression of each transporter among different human tissues and cell lines. Many transporters exhibit extremely high Gini coefficients even for common substrates, indicating considerable specialization consistent with divergent evolution. The expression profiles of SLC transporters in different cell lines behave similarly, although Gini coefficients for ABC transporters tend to be larger in cell lines than in tissues, implying selection. Transporter genes are significantly more heterogeneously expressed than the members of most non-transporter gene classes. Transcripts with the stabellest expression have a low Gini index and often differ significantly from the “housekeeping” genes commonly used for normalization in transcriptomics/qPCR studies. PCBP1 has a low Gini coefficient, is reasonably expressed, and is an excellent novel reference gene. The approach, referred to as GeneGini, provides rapid and simple characterization of expression-profile distributions and improved normalization of genome-wide expression-profiling data.

INTRODUCTION

Given that the basic genome of a differentiated organism is constant between cells (and we here ignore epigenomics), what mainly discriminates one cell type from another is its expression profile. The “surfaceome” (those proteins expressed on the cell surface) attracts our interest in particular, as it contains the transporters that determine which nutrients (and xenobiotics such as drugs) are taken up by specific cells (da Cunha et al., 2009; Palm and Thompson, 2017). Transporters are the second largest component of the membrane proteome (Almén et al., 2009), and also a (surprisingly) understudied clade (César-Razquin et al., 2015). They are classified into solute carriers (SLCs) (Colas et al., 2016; Fredriksson et al., 2008; Hediger et al., 2013; Perland and Fredriksson, 2017; Schlessinger et al., 2010; Sreedharan et al., 2011), mainly involved in uptake, and ABC transporters (ABCs), mainly involved in efflux (e.g., Chen et al., 2016; Eadie et al., 2014; Montanari and Ecker, 2015; Rees et al., 2009).

Transporters are also responsible for the uptake of pharmaceutical drugs and xenobiotics into cells, and their efflux therefrom (Colas et al., 2016; Dobson and Kell, 2008; Giacomini and Huang, 2013; Giacomini et al., 2010; Kell, 2015, 2016; Kell et al., 2011, 2013; Kell and Oliver, 2014; Lin et al., 2015; Stanley et al., 2009). This means that, to understand drug distributions, we must understand transporter distributions. In many cases, we do not know either the “natural” (O’Hagan and Kell, 2017b, 2018; Perland and Fredriksson, 2017) or the pharmaceutical drug substrates of these transporters, and one clue to this may be to understand transporters’ differential tissue distribution.

In the present work we used absolute transcription profiles acquired (via RNA sequencing) as part of the tissue atlas (Uhlen et al., 2015) and cell atlas (Thul et al., 2017). Altogether there are four main datasets, namely 409 SLCs in 59 tissue types and 56 cell lines, and 48 ABCs in the same tissue types and cell lines. Some of the SLCs do not (yet) have the official terminology (Perland and Fredriksson, 2017; Sreedharan et al., 2011), but, based on a variety of phylogenetic and other evidence, as well as their UniProt annotations, they clearly have this function, and these are noted accordingly. Similarly, some of the “ABC” families (especially family F) are probably not functionally membrane transporters, but they are nonetheless included.

The availability of extensive and high-quality transcriptomic datasets allows us to develop a series of novel analyses. They are necessarily illustrative, but by making the data available in a convenient form, we think that readers will be encouraged to make their own analyses of other aspects. In particular, the Gini index serves to highlight unusual features of the biology of a great many transcripts; we refer to this strategy of using the Gini index to analyze expression profiling data as GeneGini.
Figure 1. Overall Assessment of Variation in Gene Expression Profiles

(A) The Gini index. Many equivalent definitions are possible. In the usual form, the Gini coefficient is defined mathematically based on the Lorenz curve, which plots the proportion of the total income or wealth of a population (ordinate) that is earned cumulatively by the bottom x% of the population (see diagram) as x increases. Here “income” is the percentage of total transcripts, while the “population” is the individual transporter transcripts considered at one time. (The same general form results if the abscissa is reversed, starting with the top earners, where it takes on the appearance of the more familiar receiver-operator characteristic curve or ROC curve; Baker, 2003; Broadhurst and Kell, 2006; Linden, 2006.) The line at 45° represents uniform expression of each transcript. The Gini coefficient can then be seen as the ratio of the area that lies between the line of equality and the Lorenz curve (labeled A in the figure) to the total area under the line of equality (labeled A and B), i.e., $G = A/(A + B)$.

(B) Median and maximum expression levels (ignoring those with undetectable expression even at the median) in the 59 tissues considered.

(C) Gini coefficient for the expression of all SLCs in 59 tissues; those with Gini coefficients above 0.9 or below 0.25 are shown.

(legend continued on next page)
A preprint has been deposited at bioRxiv (O’Hagan et al., 2017).

RESULTS

Gini Index

Our first interest was to provide a convenient method for summarizing the variation in gene expression profiles in different samples (in this case different tissues and cell lines). A variety of means exist to capture variation; however, none of the more common statistical measures captures the full range well, especially including the many zeroes (undetectable expression levels). One that does is the Gini index (Ceriani and Verme, 2012; Gini, 1909, 1912) or Gini coefficient (GC). This is a non-parametric measure that is widely used in economics to describe distributions of incomes between individuals in a given group or political jurisdiction (e.g., country or region) (Kondo et al., 2012; Pickett and Wilkinson, 2015; Wilkinson and Pickett, 2009). As a summary statistic of the entire Lorenz curve (Lee, 1999) (see Figure 1), it is a statistical measure of the degree of variation represented in a set of values. It ranges between 0 (no variation) and 1 (extreme variation, in which all non-zero values are contained in one individual or example). Clearly it can be used to describe the distribution of anything else, e.g., the structural diversity in chemical libraries (Weidlich and Filippov, 2016) (modulo; O’Hagan and Kell, 2017b). It has very occasionally been used in gene expression profiling studies (Ainali et al., 2012; Jiang et al., 2016; Torre et al., 2017; Tran, 2011). However, in each of these latter cases, including a very recent and nicely done example on cancer cell susceptibility to drugs (Shaffer et al., 2017), where it varied from 0.05 to 1, the Gini index was used for choosing subsets of transcripts that differentiate rare cell types or diseases. Here we know the cell types, and the novelty of GeneGini lies in using the Gini index to assess individual genes in terms of the uniqueness of their expression levels. A more intuitive, graphical illustration is given in Figure 1A.

Variation in Expression Profiles of SLCs in Tissues

As is typical in exploratory data analysis (Tukey, 1977), we begin with the following general comments (the full datasets are given in Supplemental Information: Tables S1 and S2):

1. The variation of transporter expression levels between different tissues or cell lines is very far from being normal (Gaussian) (see Broadhurst and Kell, 2006 for methods; data not shown). The extreme here (and see below) is probably SLC01B1 (Hagenbuch and Steiger, 2013), whose expression is virtually confined to the liver alone (a fact that has been exploited effectively for drug targeting purposes [Pfefferkorn, 2013]).

2. The tissue with the maximum overall expression of transporters (SLC and/or ABCs) is the kidney ($20,950); that with the fewest is the pancreas ($1,490).

3. The SLCs with, overall, the greatest expression in total are SLC6A15 (a neutral amino acid transporter [Pramod et al., 2013]), whose activity has been implicated in depression (Kohli et al., 2011), and SLC25A3 (a mitochondrial phosphate transporter [Palmieri, 2013]), while that least expressed in toto is SLC6A5 (glycine transporter).

4. Almost every transporter ranges in its expression by over two orders of magnitude in different tissues, and several by more than three or even four orders of magnitude (see also Sreedharan et al., 2011; Winter et al., 2014).

5. The heatmap of expression levels shows a number of major co-expression clusters.

Figure S1 shows the minimum and maximum expression levels (as TPM [transcripts per million]) for each transporter, with the top 20 (maximum expressions) labeled explicitly. Open circles are those not explicitly labeled as SLC family members. Interestingly, the mitochondrial transporters (Palmieri, 2013) SLC25A3 (for phosphate) and SLC25A5 (for adenine nucleotide translocase [Clémenc¸on et al., 2013]) are among the most highly expressed, as is the non-SLC MTPCH1, which, as its name implies, is a mitochondrial carrier homologue. The co-expression of SLC25A3 and SLC25A5 is entirely logical (not shown, but see data files), as ATP synthesis and export require the transport of equimolar amounts of its substrates. Many other SLC25 (mitochondrial transporter) family members are well represented as high expressers in at least one tissue. Note that expression levels below 0.01 TPM are not shown. Figure 1B shows similar data for the median versus the maximum expression in the different tissues, which again serves to highlight the considerable heterogeneity of expression. The median of the set of median expression levels for all the SLCs was 3.19 TPM. In addition, it is not at all the case that a transporter tends to be either highly expressed or weakly expressed; although as many transporters are widely distributed, there is a considerable degree of specialization (see also Sreedharan et al., 2011).

The Gini index for the variation in (inequality of distribution of) transporters (Figure 1C) is fully consistent with this, with a significant number having an exceptionally high value (66 at 0.9 or above), not least SLC22 family members, often in the kidney (see below), and with only 23/409 SLCs having a GC below 0.25. One interpretation is that, mostly, individual transporters may be quite specialized; another is that different tissues require different amounts of specific substrates, although such large differences are thereby not easily explained in general. The median GC for this overall class of SLCs and related transporters is 0.587. A number of those with the lowest GCs are again in the SLC25 (mitochondrial transporter) family; this is not unreasonable, since every cell is likely to have mitochondria, but some family members are clearly very specialized for particular mitochondrial substrates. Thus (Figure 1D) SLC25A3 (AAC4), a particular isoform of the adenine nucleotide translocase (Palmieri, 2013), is essentially expressed only in the testes (Dolce et al., 2005) (GC = 0.965), a finding of unknown biological significance (Hamazaki et al., 2011). However, since its removal inhibits spermatogenesis (Brower et al., 2007), and thus causes infertility

(D) SLC25A31 is almost exclusively expressed in the testes (the expression levels for others being 100 times less).

(E) SLC01B1 is almost exclusively expressed in the liver (with the expression level in other tissues being 100 times lower or less).

(F) Antibody–based expression of the SLC22A12, SLC6A18, and SLC2A14 transporters in kidney, testis, and liver tissues. SLC22A12 and SLC6A18 are expressed in renal proximal tubules, whereas SLC2A14 is expressed in cells in seminiferous ducts. Image edge length is 320 μm.
SLC2A14, a glucose transporter (Mueckler and Thorens, 2013), is essentially confined to the kidney proximal tubule. Similarly, transporter, has the 15th highest GC (0.955), and its expression does not seem to hold up, and the GC again provides a convenient visual summary of large amounts of data. Thus, Figure 2B shows the full heatmap for SLC expression in tissues. Although, as stated, all the data are provided in full (Supplemental Information) to allow readers to explore them, we have marked four major clusters (zoomed in in Figures S5–S8). With the exception of a slight preponderance of families SLC 25 and 35 in cluster 3 (Figure S7) and of SLC35 in cluster 4 (Figure S8), there was no obvious clustering at the level of families. This gives weight to the idea that SLC transporters have mainly exhibited divergent evolution (Höglund et al., 2011).

SLCs in Cell Lines

Figure 3A shows the minimum non-zero versus maximum expression levels of SLCs in cell lines (Figure 3A). The trends are broadly similar, with some of the most highly expressed transporters again being SLC25A3, SLC25AS, MTCH1, and SLC3A2, although there are also differences. The overall spread seems broadly similar to those of tissues, with a preponderance of transporters having minima in the decade 1–10 TPM and maxima in the decade 20–200 TPM. In this sense, cell lines are a reasonable representation of the behavior of tissues. The number of SLCs with a GC over 0.9 is 70, while those with GCs below 0.25 is 35 (Figure 3B). These numbers and behaviors are also close to those for tissues. The median GC for SLCs in cell lines (0.595) is very close to that for tissues (0.587). We note that there may be a mixture of cell types in the tissues, and that some (or even many) transporters likely exhibit a cell-type-specific expression pattern such as SLC22A12, SLC6A18, and SLC2A14 (Figure 2). Finally (Figure 3C) we show the extensive (4,000-fold) variation in expression profiles of SLC22A4 (the ergothioneine transporter) in the different cell lines, again illustrating very substantial differences in “need” for this exogenous antioxidant (Halliwell et al., 2016) compound. Consistent with this, the cell line with the greatest expression is a skin cell line, that is normally exposed to atmospheric oxygen.

ABC Transporters in Tissues

Figure 4A shows the minimum and maximum expression levels for all 48 ABCs, many of which lack detectable expression in at least one tissue type. Again, the ranges of expression are considerable, but their expression levels tend to be slightly lower than those of the SLCs. The total numbers are small, but no family (encoded in color in Figure 5A), except possibly F, seems especially highly expressed. The overall most highly expressed ABC transporter is ABCC4. The GCs (Figure 4B) vary more than those of the SLCs, and have a median value of 0.496. Five of 48 GCs are greater than 0.9, while four are below 0.25. Several ABCs exhibit very high GCs, that (0.939) of ABCG5 being the largest; it is mainly expressed in the duodenum and the liver. Those of the F family, however, while highly expressed, also have a low GC, indicating that they tend to be among the more highly expressed in most tissues. Indeed, consistent with their being outliers, they are probably not in fact transporters (e.g., Nishimura et al., 2007).

ABC Transporters in Cell Lines

Figure 4C shows the minimum and maximum expression levels for all 48 ABCs, many of which lack detectable expression in at least one cell line. Again, the ranges of expression are considerable, and somewhat more so than those of the SLCs in tissues. No family (encoded in color in Figure 4C) seems especially highly expressed. The overall most highly expressed ABC transporter is ABCE1. The GCs (Figure 4D) are also large and vary more than those of both the SLCs and of the ABCs in tissues, with a median value of 0.692, suggesting adaptive selection for specialized purposes in the relevant cell lines. Eleven of 48 GCs are greater than 0.9, while five are below 0.25. Several ABCs exhibit very high GCs, that (0.964) of ABCG5 (a sterol transporter [Kerr et al., 2011]) again being the largest; here it is effectively expressed only in the HepG2 liver carcinoma cell line.

Overall, the median expression levels for SLCs are 3.27 and 1.26 TPM for tissues and cell lines, respectively, while those
Figure 2. Clustering of (Co-)Expression Profiles of SLC Transporters

(A) Significant correlation (in log-log space) between the expression profiles of SLC39A5 and SLC17A4 ($r^2 = 0.86$).

(B) Overall heatmap, with four major clusters highlighted.
Figure 3. Expression Profiling of Various Transporters in 56 Cell Lines
(A) Minimum and maximum expression levels (as in Figure S1 not showing those with undetectable expression) in the 56 cell lines considered.
(B) Median and maximum expression levels (ignoring those with undetectable expression even at the median) in the 56 cell lines considered.
(C) SLC22A4 expression levels (in TPM) in different cell lines.
for ABCs are 4.23 and 1.48 TPM. Thus, while many of these cell lines are cancer derived, the majority of differentially expressed genes (as transporters are) are downregulated in cancer cells (Danielsson et al., 2013). By contrast, if (as helpfully pointed out by a referee) we consider maxima, the median of the maxima in cell lines is close to double that in tissues, both for SLCs (646 versus 368 TPM) and ABCs (98 versus 48 TPM). Thus some transporters are indeed substantially overexpressed in cancer cell lines.

**Overall Analysis and Clustering of Cell Lines Based on Transporter Transcripts**

Although the data are far from being normally distributed, it is of interest to see which tissues and cell lines are most different from each other based solely on the expression profiles of their transporters; these data (normalized to unit variance) are given as a principal components plot in Figures 5A and 5B, where tissue type is encoded by color, and in the former, whether it is a tumor (gray) or not, is also encoded by a circular shape. Only a small amount of the variance is explained by the first two principal components, consistent with the high variability between tissues and cells, and scree plots are given as insets. The cell line expressing the largest total amount of transporter transcripts (11,566 TPM) in toto is BeWo (a placental carcinoma), while that expressing the fewest (5,215 TPM) is ASC TERT1 (a human telomerase-immortalized human adipose-derived mesenchymal stem cell line); the variance in transcripts that may be observed between these two cell lines is given in Figure 5C, with several of those with the greatest differences illustrated. That the total variation in transporter expression is just 2-fold shows (1) the limitation of membrane “real estate” area that partly controls membrane protein expression (Kell et al., 2015), and (2) their overall importance to the cellular economy.

**Unusually Heterogeneous Nature of Cell Transporter Expression Profiles**

**Tissues**

While the values of GC for the expression profiles of transporters between different tissues and cells tend to be unusually high, we
Figure 5. Overall Variance of SLC plus ABC Transporter Expression in Different Tissues, A, and Different Cell Lines, B

(A and B) Analyses were run in KNIME using the expression profiles of both SLCs and ABCs, each normalized to unit variance. Inserts in (A and B) represent the scree plots of percent variance explained by different principal components (PCs).

(C) Variance in transcript levels of both SLC (blue) and ABC (red) transporters in just two cell lines (BEWO and ASC/TERT1) ($r^2 = 0.50$).
have not yet quantified their differences relative to those of other genes.

From such data, the most transcribed gene over any other in cell lines is the ATP6 gene (mitochondrial ATP synthase subunit a, UniProt P00846, 42,706 TPM in HeLa cells), while that in tissues is ALB (albumin, UniProt P02768, 105,947 TPM in liver). The median of all the maxima for tissues is 46 TPM, and for cell lines 40 TPM. Obviously the first of these (ATP6 and ALB) are much larger numbers than those for any transporters (Figures 1 and 4), but the medians (see also Figure 1B) are in quite a similar range; this again illustrates the rather specialist nature of different tissue expression profiles.

The overall picture of the distribution of tissue GCs between the three classes of molecule (SLC/ABC/other) is given in Figure 6 (422 genes had very little expression at all [max = 0.25 TPM] and were ignored). Gene names are in alphabetic order, so it is clear where most of the ABCs (in blue) and SLCs (red) lie. Simply by inspection of this figure we can tell that many more “other” genes (19%) have a GC below say 0.25 than those for SLCs (9%) and ABCs (10%). In a similar vein, 33% of SLCs and 24% of ABCs have a GC exceeding 0.75, while 24% do for other genes. This latter high number is because of several clusters that are visible (and marked) in Figure 6A, specifically those for olfactory receptor proteins (over 300 genes, expressed in specific tissues, which, given their high GCs, necessarily varied for different olfactory receptor proteins) and keratin (over 150 genes, mainly in the melanoma tissues, of which 58 are KRT for keratin and 58 KRTAP for keratin-associated proteins). Note, however, that the maximum expression level for most ORs, and for 69% of the 94 KRTAP (keratin-associated protein) genes, was mainly less than 1 TPM; it is thus uncertain whether they encode detectable levels of protein. By contrast, transcriptional activators in the form of zinc-finger proteins (over 500 transcripts, 82%/97% of which had a median/maximum expression greater than 1 TPM) have very low GCs as they seem to play regulatory roles in almost all cells. Cyclins are of interest, as scripts, 82%/97% of which had a median/maximum expression encode detectable levels of protein. By contrast, transcriptional was mainly less than 1 TPM; it is thus uncertain whether they and for 69% of the 94 KRTAP (keratin-associated protein) genes, however, that the maximum expression level for most ORs, and keratin and 58 KRTAP for keratin-associated proteins). Note, therefore, that the maximum expression level for most ORs, and for 69% of the 94 KRTAP (keratin-associated protein) genes, was mainly less than 1 TPM; it is thus uncertain whether they encode detectable levels of protein. By contrast, transcriptional activators in the form of zinc-finger proteins (over 500 transcripts, 82%/97% of which had a median/maximum expression greater than 1 TPM) have very low GCs as they seem to play regulatory roles in almost all cells. Cyclins are of interest, as these should be expressed only in dividing cells. Thus CCNA1, the gene for cyclin A1, has a GC of 0.844. However, because our focus here is on transporters, we shall not pursue all these other very interesting questions here.

Genes with Low Expression Profiles as Candidate “Housekeeping” Genes

A variety of genes have previously been proposed as housekeeping or reference genes (Bustin et al., 2009; de Jonge et al., 2007; Gur-Dedeoglu et al., 2009; Hoemdl et al., 2004; Li et al., 2009; Ohl et al., 2005; Oturai et al., 2016; Silver et al., 2006; Tatsumi et al., 2008; Vandesompele et al., 2002; Wang et al., 2010; Zampieri et al., 2010).

However, the expression of most so-called housekeeping genes (that are at least expressed in all tissues) actually varies quite widely between tissues (e.g., de Jonge et al., 2007; Eisenberg and Levanon, 2003; Lee et al., 2002; Robinson and Oshlack, 2010); indeed they are sufficiently different that they can be used to classify different tissues (Hsiao et al., 2001)! Here, the housekeeping genes with the lowest GCs, hence those possibly best for normalizing transcriptome or proteome experiment, are FAM32A (an RNA-binding protein; GC = 0.137), ABCB7 (a mitochondrial heme/iron exporter; GC = 0.137), MRPL16 and MRPL21 (mitoribosomal proteins; GC = 0.138), and PCBP1 (an oligo-single-stranded-dC-binding protein; GC = 0.139). Clearly their ubiquitous distribution speaks to their essentiality, and it is certainly of interest that mitoribosomal proteins have such ubiquitous expression, being somewhat equivalent to the 16S rRNA genes widely used in microbial taxonomy and metagenomics. Most of the other 49 large (MRPLxx) and 30 small (MRPSxx) ribosomal protein subunits also had low GCs; others with a GC of 0.15 or below are illustrated in Figure 6B, which also serves to show that most low-Gini gene products have median expression levels in the decade 20–200 TPM (so it is not a strange low-expression artifact).

We note that Eisenberg and Levanon (2013) provide a list of candidate housekeeping genes based on earlier RNA sequencing data. This provides a valuable benchmark for comparison with our approach. However, their list (see http://www.tau.ac.il/~elleis/HKG/HKG_genes.txt) consists of no fewer than 3,804 genes (out of the ~25,000 human genes), but provides no quantification of either how good they are as housekeeping/referene genes or of their typical expression levels. Finding the best 6 or 7 out of such an unranked list of 3,804 is a combinatorial problem that would require testing 4.1018 or 2.1021 combinations, respectively. By contrast we provide both the rank order (and its justification via the Gini index) and the transcription level. Secondly, the paper itself (Eisenberg and Levanon, 2013) used only 16 (not, as here, 59) tissues, and no cell lines. Thirdly, the paper does contain a Table of eleven “genes proposed for calibration”, representing (on an unstated basis) “a short list of highly uniform and strongly expressed genes that may be used for calibration in future experimental settings”; Table S3 lists these, together with their correct names, UniProt ID, and (from our data) Gini index and median tissue expression levels.

It is rather obvious (Table S3) that the choices in this Table are far poorer than those we suggest in terms of both GC (only one has a GC below 0.15 [for tissues we show 23] Figure 6B) and expression level (e.g., PCBP1 has a GC of 0.139 and an expression level of 209 TPM in tissues).

Indeed, the GCs of other gene products commonly used by experimental biologists to normalize expression profiles were often considerably larger (Table S4), although the more recently proposed CTBP1 (C-terminal-binding protein 1, UniProt Q13363; 0.204) and GOLGA1 (Golgin subfamily A member 1, UniProt Q92805; 0.189) (Lee et al., 2007) both seem like much better choices. However, the lowest GCs in tissues are FAM32A, ABCB7, MRPL21, and PCBP1 (GC = 0.137–0.139), while the lowest three in cell lines are SF3B2, NFX1, and RBM45 (GC = 0.115–0.122). PCBP1 is both reasonably highly expressed and has a low GC in both tissues (0.139) and cell lines (0.135), and is an excellent novel housekeeping gene. While reference genes are often chosen to be stably expressed across variants of the same cell type rather than across different cells, our very low GC between cell types suggests that the GC is indeed a novel and effective way of identifying very useful housekeeping or reference genes in expression profiling studies.

While there was no relationship between the GC and the maximum expression (not shown), there was an interesting inverse relationship between the GC and the median expression level over all genes (Figure 6C), where the correlation coefficient was 0.62. Clearly the exact correlation is also likely to depend on the value of the GC, where at higher levels the Lorenz curve...
(Figure 1) can become highly nonlinear. The overall distribution of GCs for the three classes of protein (SLC/ABC/other) is given in Figure 6D. Finally, because it was one of the gene products with the lowest GC, as well as having a reasonable expression level (median over 100 TPM), in both tissues and cell lines, we show the tissue expression profile of PCBP1 (an intronless gene; Makeyev et al., 1999) in Figure 6E; the overall variation of the great majority of these transcripts is within a 2-fold range. We
also illustrate its distribution in several tissues in Figure 6F. This makes a very strong case for it being a highly useful reference or housekeeping gene.

**Cell Lines**

The overall data are broadly similar for cell lines (Figure 7A, although the expression of zinc fingers is less homogeneous.
expression level ($r^2 = 0.67$) (Figure 7C). Overall, we find that there is again a correlation between the Gini index and median expression among the Gini index here ranged from as low as 0.11 to as high as 0.98, reflecting in the latter case virtually unique expression in a particular tissue. In many cases, the biology underpinning this is quite opaque, but the purpose of data-driven studies is to generate rather than to test hypotheses (Kell and Oliver, 2004). We also recognize here that we have paid relatively little attention to the distribution of transporters within different tissues and their potential cell-type-specific distribution within an organ (e.g., Bahar Halpern et al., 2017), where they presumably account for the very striking intra-organ distributions of drugs (e.g., Römpf et al., 2011); that will have to be a subject for further work.

A second chief area of interest is the distribution of transporters between different tissues. A detailed analysis showed that they tended to have significantly higher GCs than did other gene families. This illustrates the point that despite the fact that their substrates are almost uniformly available via the bloodstream, and biochemistry textbooks and wallcharts largely show this, they clearly use substrates differentially (ergothioneine and the SLC22A4 transporter being a nice example; Gründemann et al., 2005). It also implies strongly that in many cases we do not in fact know the natural substrates, many of which are clearly exogenous (O’Hagan and Kell, 2017b).

The third main recognition is that the Gini index provides a particularly useful, convenient, non-parametric, and intelligible means of identifying those genes whose expression profile varies least across a series of cells or tissues, thus providing a novel and convenient strategy for the identification of those reference or housekeeping genes best used as genes against which to normalize other expression profiles in a variety of studies. We have here highlighted quite a number that have not previously been so identified.

Overall, we consider that assessing the Gini index for the distribution of particular transporters and other proteins between different cells has much to offer the development of novel biology; it should prove a highly useful addition to the armory of both the systems biologist and the data analyst.

STAR+METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- CONTACT FOR REAGENT AND RESOURCE SHARING
- METHOD DETAILS
  - Gini Index
  - Minimum and Maximum Expression Profiles
- QUANTIFICATION AND STATISTICAL ANALYSIS
- DATA AND SOFTWARE AVAILABILITY

SUPPLEMENTAL INFORMATION

Supplemental Information includes eight figures and four tables and can be found with this article online at https://doi.org/10.1016/j.cels.2018.01.003.

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AUTHOR CONTRIBUTIONS

The project was initiated following an initial discussion between E.L. and D.B.K. at a scientific conference. D.B.K. highlighted the utility of the GC, and produced many of the visualisations. S.O. adapted the Gini method and performed most of the analyses that were done using KNIME. M.W.M. did the principal-component analyses, while P.J.D. contributed in particular to the analysis of the housekeeping genes. E.L. contributed the original data and images, and multiple insights thereto. All authors contributed to the writing and approval of the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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**STAR METHODS**

**KEY RESOURCES TABLE**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Software and Algorithms** | Gini coefficient | https://CRAN.R-project.org/package=ineq |
| **RESOURCE: Cell Atlas, cell line RNA-seq data** | Human Protein Atlas | https://www.proteinatlas.org/download/rna_cellline.tsv.zip |
| **RESOURCE: Tissue Atlas, tissue RNA-seq data** | Human Protein Atlas | https://www.proteinatlas.org/download/rna_tissue.tsv.zip |

**CONTACT FOR REAGENT AND RESOURCE SHARING**

Further information and requests for resources should be directed to and will be fulfilled by the Lead Contact, Douglas B. Kell (dbk@manchester.ac.uk).

**METHOD DETAILS**

The expression profile data are not new; the means by which they were obtained is described elsewhere (Thul et al., 2017; Uhlén et al., 2015). mRNA sequencing was performed on Illumina HiSeq2000 and 2,500 platforms (Illumina, San Diego, CA, USA) using the standard Illumina RNA-seq protocol with a read length of 2x100 bases. Transcript abundance estimation was performed using Kallisto (Bray et al., 2016) v0.42.4. For each gene, we report the abundance in 'Transcripts Per Million' (TPM) as the sum of the TPM values of all its protein-coding transcripts. For each cell line and tissue type, the average TPM values for replicate samples were used as abundance score. Thus each transcript level does represent an absolute value, but it is then normalised to the total expression in the particular sample. The data were extracted and extended in the form of Microsoft Excel sheets (Raw SLC and ABC data in Tables S1 and S2).

Most of the analyses are self-explanatory, but are noted below. As in many of our cheminformatics analyses (e.g. (O’Hagan and Kell, 2017a; O’Hagan et al., 2015)) we used the freely available KNIME software environment (Berthold et al., 2008; O’Hagan and Kell, 2015; O’Hagan et al., 2015) (http://knime.org/), with visualisation often provided via the Tibco Spotfire software (Perkin-Elmer Informatics).

**Gini Index**

The Gini Index was calculated using the ineq package (Achim Zeileis (2014). ineq: Measuring Inequality, Concentration, and Poverty. R package version 0.2-13. https://CRAN.R-project.org/package=ineq) in R (https://www.R-project.org/). These calculations were incorporated into KNIME via KNIME’s R integration R Snippet node. The Rank Correlation used was Spearman’s rho, using the KNIME Rank Correlation node.

**Minimum and Maximum Expression Profiles**

These and the other similar analyses were done using the functions contained in MS-Excel.

**Immunohistochemistry**

Immunohistochemical (IHC) images detailing protein expression patterns in 48 different normal tissues and 20 common cancer types are from the Human Protein Atlas database (www.proteinatlas.org). Tissue microarrays, immunostaining and image evaluation was performed as previously described (Uhlén et al., 2015). Briefly, 1mm duplicate cores were used for immunostaining using the following antibodies: HPA024575 for SLC22A12, HPA011885 for SLC6A18, HPA006539 for SLC2A14 (all from the Human Protein Atlas) and CAB037113 for PCBP1 (R1455 from Sigma-Aldrich). The immunostaining intensity and pattern was manually evaluated and scored by certified pathologists.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

For each cell line and tissue type, the average TPM values for replicate samples were used as abundance score.

**DATA AND SOFTWARE AVAILABILITY**

The data on which we base our analyses are all available online at https://www.proteinatlas.org/about/download (and see Key Resources Table).
Supplemental Information

GeneGini: Assessment via the Gini Coefficient of Reference “Housekeeping” Genes and Diverse Human Transporter Expression Profiles

Steve O'Hagan, Marina Wright Muelas, Philip J. Day, Emma Lundberg, and Douglas B. Kell
SUPPLEMENTARY INFORMATION

The extra subsetted data that we give in the Supplementary Information are as follows:

**Supplementary Table S1.** Related to Fig 1. Expression profiles of the SLC transporters. Separate XL File: Supplementary Table S1 SLC_transporters_RNA_data_HPA.xlsx

**Supplementary Table S2.** Related to Fig 4. Expression profiles of the ABC transporters. Separate XL File: Supplementary Table S2 ABC_transporters_RNA_data_HPA.xlsx

**Supplementary Table S3.** Related to STAR Methods. A previously proposed set of useful reference genes, annotated here with their correct names and Uniprot IDs, together with their median expression levels and Gini indices in tissues as determined in this work.

**Supplementary Table S4.** Related to STAR Methods. Some genes that have previously been proposed as housekeeping or reference genes.

**Supplementary Figures.**

**S1.** (Relates to Fig 1.) Expression profiling of various SLC transporters in 59 tissues. Minimum and maximum expression levels of various SLCs in the 59 tissues considered (those with undetectable expression (i.e. <0.01 TPM, coded as zero) are not shown).

**S2.** (Relates to Fig 1.) Expression profiling of various SLC transporters in 59 tissues. The expression level of SLC35A4 is relatively homogeneous, with ¾ of all tissues within a factor two.

**S3.** (Relates to Fig 1.) Expression profiling of various SLC transporters in 59 tissues. The expression levels of SLC35F2 vary much more considerably, by a range of ~200 in these 59 tissue types.

**S4.** (Relates to Fig 1.) Expression profiling of various SLC transporters in 59 tissues. Expression profile of the transcripts for SLC22A4.

**S5.** (Relates to Fig 2.) Zoomed-in version of cluster 1 of Figure 2.

**S6.** (Relates to Fig 2.) Zoomed-in version of cluster 2 of Figure 2.

**S7.** (Relates to Fig 2.) Zoomed-in version of cluster 3 of Figure 2.

**S8.** (Relates to Fig 2.) Zoomed-in version of cluster 4 of Figure 2.
| Gene name | Protein name                                           | Uniprot ID | Gini index in tissues | Median expression level (TPM) |
|-----------|--------------------------------------------------------|------------|-----------------------|-----------------------------|
| C1orf43   | Chromosome 1 open reading frame 43                     | Q9BWL3     | 0.204                 | 137                         |
| CHMP2A    | Charged multivesicular body protein 2A                 | O43633     | 0.141                 | 126                         |
| EMC7      | ER membrane protein complex subunit 7                 | Q9NPA0     | 0.210                 | 69                          |
| GPI       | Glucose-6-phosphate isomerase                          | P06744     | 0.259                 | 137                         |
| PSMB2     | Proteome subunit beta type 2                           | P49721     | 0.186                 | 32                          |
| PSMB4     | Proteome subunit beta type 4                           | P28070     | 0.200                 | 209                         |
| RAB7A     | RAS-related protein 7A                                 | P51149     | 0.171                 | 167                         |
| REEP5     | Receptor expression-enhancing protein 5                | Q00765     | 0.315                 | 65                          |
| SNRPD3    | Small nuclear ribonucleoprotein Sm D3                  | P62318     | 0.192                 | 55                          |
| VCP       | Transitional endoplasmic reticulum ATPase (originally valosin containing protein) | P55072 | 0.198 | 48            |
| VPS29     | Vacuolar protein sorting associated protein 29         | Q9UBQ0     | 0.146                 | 74                          |

**Supplementary Table S3.** A previously proposed set of useful reference genes, annotated here with their correct names and Uniprot IDs, together with their median expression levels and Gini indices in tissues as determined in this work.
| Gene   | Protein                                      | Uniprot ID | Gini index |
|--------|----------------------------------------------|------------|------------|
| GAPDH  | Glyceraldehyde 3-phosphate dehydrogenase     | P04406     | 0.344      |
| LDHA   | Lactate dehydrogenase subunit A              | P00338     | 0.32       |
| SDHA   | Succinate dehydrogenase subunit A            | P31040     | 0.308      |
| HRPT1  | Hypoxanthine phosphoribosyl transferase 1    | P00492     | 0.277      |
| HBS1L  | HBS1-like protein                            | Q9Y450     | 0.184      |
| OAZ1   | Ornithine decarboxylase antizyme 1           | P54368     | 0.202      |
| PPIA1  | Peptidyl-prolyl cis-trans isomerase          | P62937     | 0.24       |
| AHSP   | Alpha-haemoglobin stabilising protein        | Q9NZD4     | 0.97       |
| B2M    | β2-microglobulin                             | P61769     | 0.349      |
| ACTB   | β-actin                                      | P60709     | 0.291      |
| HMBS   | Porphobilinogen deaminase                    | P08397     | 0.303      |
| UBC    | Polyubiquitin C                              | P0CG48     | 0.183      |
| POLR2F | DNA-directed RNA polymerases I, II, and III subunit RPABC2 | P61218 | 0.235 |
| GUSB   | β-glucuronidase                              | P08236     | 0.25       |
| TBP    | TATA-box binding protein                     | P20226     | 0.22       |
| YWHAZ  | 14-3-3 protein zeta/delta                    | P63104     | 0.255      |

**Supplementary Table S4.** Some genes that have previously been proposed as housekeeping or reference genes.
Supplementary Fig S1 (Relates to Fig 1.) Expression profiling of various SLC transporters in 59 tissues. Minimum and maximum expression levels of various SLCs in the 59 tissues considered (those with undetectable expression (i.e. <0.01 TPM, coded as zero) are not shown).
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Supplementary Fig S8. (Relates to Fig 2.) Zoomed-in version of cluster 4 of Figure 2.