Identification of putative calorie restriction mimetics using mammalian gene expression profiles

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Obesity is a risk factor for cardiovascular diseases, diabetes and cancer. In theory, the obesity problem could be solved by the adherence to a calorie-restricted diet, but that is not generally achieved in practice. An alternative is a pharmacological approach, using compounds that trigger the same metabolic changes associated with calorie restriction. Here, I expand in the pharmacological direction by identifying compounds that induce liver gene signature profiles that mimic those induced by calorie restriction. Using gene expression profiles from mice and rat, I identify corticosteroids, PPAR agonists and some antibacterial/antifungal as candidate compounds mimicking the response to calorie restriction in the liver gene signatures.

1. Background

Obesity results from an excess calorie intake relative to expenditure [1]. It seems reasonable that calorie restriction would be sufficient to tackle this problem. However, for a number of reasons, long-term calorie restriction is hard to sustain [2]. A valid alternative is to search for pharmacological agents that could achieve weight loss while maintaining a regular or less restricted diet [3,4]. Some advances have been made in this direction, with the development of PPARα and PPARγ agonists that stimulate oxidation of fatty acids [5–7].

Drug repurposing is a potential strategy to identify additional compounds to mimic calorie restriction [8]. A popular approach has been to search for compounds that match a specified gene expression profile, based on the in vitro cell culture response to a large library of compounds [9]. This approach has led to the identification of putative calorie restriction mimetics [10]. Given the proven success of the gene expression profile matching, it is desirable to develop a similar methodology to tackle disease phenotypes that are mainly manifested in mammalian tissues.

Here, I propose a gene signature methodology to identify candidate compounds that trigger a target gene signature profile. As a bait, I use gene expression profiles from the liver of mice subject to calorie restriction, downloaded from Gene Expression Omnibus. As a probe, I use gene expression profiles from the liver of rat exposed to a large collection of compounds from a toxicology study [11]. I identified corticosteroids, PPAR agonists and some antibacterial/antifungal agents as candidate compounds to mimic calorie restriction.

2. Methods

2.1. Gene signatures

All gene signatures were obtained from public repositories or literature reports. The source and gene list of each signature is reported in electronic supplementary material, table S1.
2.2. Gene expression profiles

The gene expression profiles were downloaded from Gene Expression Omnibus. In all cases, gene expression was quantified using microarrays and the RMA signal (in log2 scale) was used as the gene expression readout.

2.3. Gene set enrichment analysis

The significant induction/repression of a given gene signature on a given sample was quantified using gene set enrichment analysis (GSEA) [12], as previously described [13]. GSEA results in a positive and a negative score quantifying the induction or repression of the gene signature, together with their associated statistical significance (here 100 000 permutations of the gene assignment to probes). A sample was defined positive (red) for a signature whenever it manifested a significant positive score (statistical significance $\leq 0.05$), negative (blue) whenever it manifested a significant negative score (statistical significance $\leq 0.05$) and no significant change (black) otherwise.

2.4. Similarity statistics

Given the signature scores for two samples, at the red/black/blue level, the similarity was calculated using the Pearson correlation coefficient (PCC). Given two groups of samples associated with two different perturbations, the PCC was calculated between all possible pairings between the two groups and the values above the 5%, 50% (median) and 95% of all calculated PCCs were stored for further analysis.

2.5. Positive hits

Given a bait gene signature scores, a probe gene signature score was deemed a positive hit if the 5% value was positive. For each compound tested in our probe dataset, there were several conditions using different doses and treatment durations. Each of these conditions was processed as an independent test. A compound was deemed a hit if it has a significant enrichment of positives across the conditions where it was tested, given the total number of conditions tested and the total number of conditions scoring positive in the whole dataset, using a hypergeometric distribution (electronic supplementary material, table S2).

3. Results

3.1. Gene signature selection

Here, I will use liver gene expression profiles as a way to identify compounds mimicking calorie restriction. These profiles report the expression of more than 20 000 genes as measured using microarrays or RNA sequencing. This large number of variables could carry as a consequence overfitting towards any given dataset, making the biological interpretation of any result difficult. There are multiple approaches to reduce the number of variables. The standard choice is unsupervised methods such as principal component analysis or its nonlinear generalization using auto-encoders. However, these approaches will be biased for the datasets used to determine the principal components. The analysis of gene signatures provides an alternative supervised approach, with the additional advantage of improving on the biological interpretation.

Here, I will adopt a gene signature approach. Gene signatures are gene lists based on pathway annotations, genes changing their expression under pre-defined perturbations, or any biological mean associating a group of genes. Ironically, the number of gene signatures in repositories like the molecular signatures database (MSigDB) [14] is getting close to 20 000, bringing us back to the overfitting problem. To tackle this issue, I will limit the analysis to a restricted set of signatures with relevance to calorie restriction (electronic supplementary material, table S1).

The list starts with two fundamental processes of organ homeostasis, cell proliferation and tissue remodelling, followed by gene signatures associated with central metabolism, fatty acid metabolism, cholesterol metabolism and one-carbon metabolism. I will use another subset of signatures to interrogate the potential activation of relevant transcriptional programs. This includes MYC targets as an additional readout for cell proliferation, HIF1α targets for the hypoxic response, NRF2 targets for inflammation, ATF4 targets for amino acid stress response, NFR2 targets for oxidative stress response, HSFI targets for proteotoxic stress response and TFE3B targets for an autophagic response. The list of signatures ends with variants of cell death, including apoptosis, necrotic cell death and ferroptosis, plus DNA repair as a readout for DNA damage response.

3.2. Gene signature validation

The signatures of cell proliferation and tissue remodelling have been tested in the context of human tumours [13]. The metabolic signatures are based on established gene annotations of metabolic pathways. The transcription factor targets’ signatures have been developed for the most part from experiments with cells in culture. Therefore, it is important to determine whether they reflect their associated biology in whole tissues. To tackle this problem, I searched the Gene Expression Omnibus database for gene expression profiles associated with the relevant biological conditions.

I identified a dataset reporting liver gene expression profiles of mice in a hypoxic chamber (6–8% oxygen) for 2 h (GSE17880 [15]). In this dataset, there is a significant induction of the gene signature of HIF1α targets in the liver of mice under hypoxic conditions relative to controls (figure 1a). Interestingly, there is some variability with respect to the metabolic signatures in the group of mice exposed to hypoxia. Yet, in both groups, there is a consistent induction of the HIF1α targets signature.

I identified another dataset reporting liver gene expression profiles of mice injected with scrambled or liver-specific siRNA against Keap1, the gene encoding the main negative regulator of NRF2 (GSE80956 [16]). I found a clear induction of the NRF2 targets gene signature in the mice treated with the liver-specific Keap1 siRNA (figure 1b).

I identified a third dataset where the human TFE3 was injected into the liver of mice using an adenovirus transduction system (GSE35015 [17]). Here again there is the matching induction of the TFE3 targets gene signature in mice injected with human TFE3 expressing adenovirus relative to non-injected controls (figure 1c). In this context, there is also induction of the HIF1α and NRF2 targets.
signatures. This could be biologically relevant. TFEB induces an autophagy program that can lead to mitophagy and the associated reduction in mitochondrial content. The latter could in turn trigger a hypoxic and oxidative stress response, leading to the induction of the HIF1α and NRF2 targets signatures. Whether that is actually the case is beyond the scope of this work.

I also took advantage of the exogenous TFEB expression dataset containing liver gene expression profiles of Pparaα−/− mice. In the liver of these mice, there is a downregulation of the PPARα targets signature as would be expected (figure 1d). In addition, the liver of Pparaα−/− mice exhibits downregulation of the fatty acids β-oxidation gene signature, one of the main processes induced by PPARα.

Unfortunately, I could not find any liver gene expression profile suitable to test the ATF4 targets signatures. In this case, I rely on gene expression profiles of the human breast cancer cell line MCF7 cultured under amino acid deprivation (GSE62673 [18]). There is a consistent induction of the ATF4 targets signature under the deprivation of each amino acid relative to cells cultured in complete medium (figure 1e). Serine synthesis and mitochondrial one-carbon metabolism, two pathways under the control of Atf4 [19–21], were also induced upon amino acids deprivation (figure 1d).

3.3. Validation across independent experiments

In a second validation, I evaluated the consistency of the signature profiles across two independent experiments testing the same perturbation in the same organism. To this end, I identified two liver gene expression profiles (GSE21329 [22] and GSE57815 [11]) of rats treated with the PPAR agonists pioglitazone and troglitazone. After computing the significance scores for each gene signature on each liver, I calculated the Pearson correlation coefficient between the signature profiles of different experiments for each compound. For pioglitazone, I obtained a median correlation of 0.37 with confidence interval [0.16, 0.49] and for troglitazone a median of 0.32 with confidence interval [0.20, 0.44]. The concordance between the signature profiles is also visually observed (figure 2a and b).

One could argue that while these correlations have confidence intervals on the positive side they are not too high. However, we should bear in mind that there are differences in the protocols used by each independent experiment. One study (GSE59927) used the common rat, higher compound doses (pioglitazone 1500 mg kg−1 and troglitazone 1200 mg kg−1) and short treatment time (3 days). The other study (GSE21329) used Zucker obese rats, lower doses (pioglitazone 10 mg kg−1 and...
3.4. Gene signature profile of calorie restriction

Now I switch the attention to the bait, the gene signature profile of calorie restriction. To this end, I selected datasets from Gene Expression Omnibus reporting gene expression profiles of mice subjected to dietary restriction (GSE51885 [23]), or calorie restriction of different magnitude and duration (GSE50789 [24] and GSE40936 [25]).

The gene signature profiles exhibit some similarities across the different experiments and across mouse strains (figure 3a). There is a consistent induction of the gene signatures associated with central metabolic pathways (glycolysis, pentose phosphate pathway, TCA cycle and oxidative phosphorylation). However, there is some variability for the pentose phosphate pathway, TCA cycle and oxidative phosphorylation. The GSE51885 dataset, representing dietary restriction with a 25% reduction of all nutrients present in the control diet, exhibits similar gene signatures to the GSE50789 dataset, representing a 25% calorie restriction. This suggests that the calorie restriction component of the dietary restriction is responsible for the observed gene signatures.

Summarizing, in the context of mild calorie restriction (approx. 25%), the liver exhibits an induction of gene signatures associated with central metabolism, HIF1α targets and TFEB targets. I note that these changes are consistent with the underlying physiology. It has been shown that autophagy is a modulator of the impact of calorie restriction on longevity [26–28]. The induction of HIF1α targets is most likely a downstream effect, as seen from the injection of TFEB expressing adenovirus into the liver of mice (figure 1c).

3.5. Compounds matching the calorie restriction gene signature

Having identified the gene signature pattern induced by calorie restriction, we can use it as a bait to uncover candidate compounds to mimic calorie restriction. To this end, we need access to liver gene expression profiles following treatment with several compounds. Fortunately, a large-scale
toxicology study in rats has profiled 194 compounds (including different vehicles) at different durations (0.25 to 7 days) and reported the corresponding liver gene expression (GSE57815 [11]). I will use the gene expression profiles in this dataset as probes to identify compounds that trigger a gene signature response similar to what observed for calorie restriction and antibacterial/antifungal agents provides two new classes of putative agents for the stimulation of a calorie restriction transcriptional program.

A previous study has reported similarities between the gene expression profile induced by calorie restriction and the PPARα/γ agonist rosiglitazone, given further confidence in the robustness of the methodology. PPARα/γ agonists have been developed for the management of metabolic syndrome [29]. The identification of corticosteroids and antibacterial/antifungal agents provides two new classes of putative agents for the stimulation of a calorie restriction transcriptional program.

Table 1 reports the best hits for each dataset and mouse strain. The full list of hits is reported in the electronic supplementary material, table S2. There are three major classes of compounds identified in two or three conditions (dataset/strain): corticosteroids, PPARα/γ agonists and some antibacterial agents. The identification of PPARα/γ agonists is a validation of the methodology. PPARα/γ agonists have been developed for the management of metabolic syndrome [29]. The identification of corticosteroids and antibacterial/antifungal agents provides two new classes of putative agents for the stimulation of a calorie restriction transcriptional program.
| dataset | GSE51885 | GSE50789 | GSE50789 | GSE50789 | GSE50789 | GSE50789 | GSE50789 | GSE40936 |
|---------|----------|----------|----------|----------|----------|----------|----------|----------|
| strain  | C57BL/6  | B6C3F1/J | C57BL/6  | C57BL/6  | C57BL/6  | C57BL/6  | C57BL/6  | C57BL/6  |
| gender  | male     | male     | male     | male     | male     | male     | male     | male     |
| age     | 10 weeks | 8 weeks  | 8 weeks  | 8 weeks  | 8 weeks  | 8 weeks  | 8 weeks  | 27 weeks |
| calorie restriction | 25% less food | 25% less calories | 25% less calories | 25% less calories | 25% then 42% less calories | 25% then 42% less calories | 40% less calories |
| duration | 3 weeks  | 14 weeks | 14 weeks | 14 weeks | 14 weeks | 14 weeks | 14 weeks | 30 weeks |
| corticosteroids | cortisone, betamethasone, dexamethasone, fluocinolone, acetonide, hydrocortisone | cortisone, hydrocortisone | fluocinolone, acetonide | fluocinolone acetonide | cortisone, betamethasone, dexamethasone, fluocinolone, acetonide | |
| anti-inflammatory | | piclamilast, balsalazide | balsalazide, benzethonium chloride, nystatin | piclamilast, benzethonium chloride, gentamicin, nystatin | |
| antibacterial/ | enoxacin | enoxacin, nystatin | enoxacin | benzethonium chloride, nystatin | |
| antifungal | | | | | |
| PPARγ agonists | bezafibrate, fenofibrate, gemfibrozil, pioglitazone | bezafibrate, fenofibrate, gemfibrozil, pioglitazone | bezafibrate, fenofibrate, gemfibrozil, pioglitazone | bezafibrate, fenofibrate, pioglitazone |
| oestrogens/ | diethylstilbestrol, estiol, norethindrone, acetate, norethindrone, progesterone | | | |
| progestogens | | | | | |
| anti-androgen | spironolactone, Bis[2-ethylhexyl]phthalate finasteride | spironolactone, Bis[2-ethylhexyl]phthalate finasteride | | |
| other | sodium aminate, 2-acetylaminofluorene, itraconazole | gefitinib, cyclopropane, carboxylic acid | artemisinin, sotalol | aflatoxin B1, sotalol | clofibrate acid, clofibrate, itraconazole, nafenopin, naphthoic acid | | | |
to the identification of this class of compounds [30]. The gene signature similarity between calorie restriction and PPARα/γ agonist treatment can be visually inspected in figure 3a and b. As calorie restriction, PPARα/γ agonists induce gene signatures of central metabolism (TCA cycle and oxidative phosphorylation), fatty acid metabolism and PPARα targets signatures. However, as a difference with calorie restriction, the PPARα/γ agonists do not induce the gene signatures of glycolysis and the pentose phosphate pathway and only Pioglitazone induces the TFEB targets signature.

Among the compounds scoring the highest, I found the synthetic corticosteroids dexamethasone and fluocinolone acetonide, often used as anti-inflammatory agents. The gene signatures associated with these compounds include the induction of HIF1α targets and partially the induction of TFEB targets (figure 3c). They also induce some of the signatures of central metabolism (TCA cycle and oxidative phosphorylation) but fail to induce the signatures of glycolysis and the pentose phosphate pathway, as seen for calorie restriction (figure 3a).

The other new class of candidate calorie restriction mimics is antibacterial/antifungal agents (figure 3d). They induce the gene signatures associated with the TFEB and HIF1α targets, but they tend to induce a downregulation of the gene signature associated with central metabolism. I note that although these compounds are all classified as antibacterial/antifungal agents they have different mechanisms of action. Nystatin is an interesting candidate since it binds to phospholipid membranes and this binding is enhanced by the addition of cholesterol [31]. In fact, nystatin is frequently used as a tool to lower cholesterol and this activity leads to the induction of autophagy [32]. In agreement with these known facts, nystatin induces the gene signature associated with cholesterol synthesis (figure 3d).

4. Discussion

This work demonstrates the use of liver gene expression signatures from mammalian model organisms subjected to calorie restrictions as baits to identify compounds that induce a similar transcriptional response. In turn, we can use as probes the liver gene expression signatures induced by treatment with a library of compounds. Here, I have used data from the public domain as a proof of concept and identified corticosteroids and some antibacterial/antifungal agents as new candidate compounds to mimic calorie restriction.

The identification of PPAR agonists, as calorie restriction mimics based on the liver gene expression profiles, resonates with a recent study reporting the same conclusion, but based on gene expression profiles for adipose, skeletal muscle, heart and brain tissues [33]. Therefore, at least for this class of compounds, there is evidence that the similarity with the calorie restriction response extends to multiple tissues.

In the case of corticosteroids, the list of candidate compounds contains synthetic molecules like dexamethasone and fluocinolone acetonide and endogenous molecules like cortisone. Cortisone is produced from cholesterol in the adrenal gland and then converted to cortisol in the adrenal gland, the liver and other tissues. Cortisone can be supplemented as well. Calorie restriction increases circulating cortisol levels [34], indicating a causal link for the similarity of the liver expression profiles induced by calorie restriction and corticosteroids. The conversion of cortisone to cortisol in the liver is reduced in obese individuals resulting in lower circulating levels of cortisol [35]. However, the pathophysiology of cortisol in the context of obesity is a complex matter [36], making it difficult to anticipate whether the administration of synthetic corticosteroids would provide any benefit or worsen the disease symptoms. Furthermore, high levels of cortisone are associated with cognitive impairment [37]. This should be further investigated.

The identified antibacterial/antifungal agents provide additional candidate compounds that should be followed as well. The specific compounds analysed here were constrained to what was included in the toxicology screen. They were not optimized by any means to induce a calorie restriction response in the liver. The identification of nystatin, a compound that lowers cholesterol and induces autophagy, highlights a common feature of calorie restriction, the induction of an autophagy gene signature in the liver. Whether that is the most relevant or a leading feature remains to be determined.

There is plenty of room for improvement. One can expand the compounds tested to a library that is more relevant than the one used in the toxicology study. One can fine tune the gene signature list to include other features deemed relevant by human experts or artificial intelligence programs. Finally, the same approach can be deployed to tackle other diseases.

5. Conclusion

Corticosteroids and some antibacterial/antifungal agents are new candidate compounds to induce a liver transcriptional response similar to calorie restriction.

Data accessibility. All the gene expression data analysed in this work are available from Gene Expression Omnibus.

Authors’ contributions. A.V. conceived the work, conducted the analyses and wrote the manuscript.

Competing interests. The author declares no competing interests.

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