siRNA Knockdown of Ribosomal Protein Gene RPL19 Abrogates the Aggressive Phenotype of Human Prostate Cancer

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Abstract

We provide novel functional data that posttranscriptional silencing of gene RPL19 using RNAi not only abrogates the malignant phenotype of PC-3M prostate cancer cells but is selective with respect to transcription and translation of other genes. Reducing RPL19 transcription modulates a subset of genes, evidenced by gene expression array analysis and Western blotting, but does not compromise cell proliferation or apoptosis in-vitro. However, growth of xenografted tumors containing the knocked-down RPL19 in-vivo is significantly reduced. Analysis of the modulated genes reveals induction of the non-malignant phenotype principally to involve perturbation of networks of transcription factors and cellular adhesion genes. The data provide evidence that extra-ribosomal regulatory functions of RPL19, beyond protein synthesis, are critical regulators of cellular phenotype. Targeting key members of affected networks identified by gene expression analysis raises the possibility of therapeutically stabilizing a benign phenotype generated by modulating the expression of an individual gene and thereafter constraining a malignant phenotype while leaving non-malignant tissues unaffected.

Introduction

Ribosomal proteins (RPs) comprise a complex super-family of proteins [1] highly conserved throughout evolution, indicating their functional importance to living organisms [2]. This assertion is supported by the number of RP pseudogenes and gene duplications together with shared regions of identity between homologous proteins in prokaryotes and eukaryotes [3]. Eukaryotic ribosomes contain approximately 80 RPs together with four ribosomal RNAs (rRNA) and require some 150 non-ribosomal factors to become organized into their constituent small (40S) and large (60S) subunits [4]. Initially considered to be involved only in protein synthesis, certain RPs are recognized as pleiotropic and to mediate a variety of extra-ribosomal regulatory functions [5,6]. Such RPs, include L5 [7], L11 [8], L13 [9] and S7 [10]. In zebrafish (Danio rerio) a powerful role for RPs as tumor-suppressors has been demonstrated whereby mutation or suppression in any of several RP genes impairs control of p53, thus promoting malignancy [11,12]. Recently, the concept of “ribosomalopathy” has become established whereby mutation of a particular RP is pathogenic for a specific disease [13]. Approximately 25% of cases of Diamond-Blackfan anemia are caused by mutation of ribosomal protein gene RPS19 while in another 20%, mutations occur in other ribosomal protein genes [14]. Currently, some 77 individual RPS19 mutations have been described [15]. In addition, haploinsufficiency for ribosomal proteins has been shown, in some cases, to be an underlying cause for Diamond Blackfan anemia [16].

Presently, mechanisms relating mutations in RP genes to cancer remain unknown [17]. For the proximal long arm region of chromosome 17 where the RPL19 gene is located (17q), major cancer-specific changes have been described. These include amplifications and copy number changes, particularly those of the region that include oncogene ERBB2, formation of isochromosome 17q, duplications, deletions, mutations and other genomic rearrangements. Previously [18], we identified enhanced expression of RPL19 mRNA in prostate cell-lines and tissues to correlate with an aggressive malignant phenotype. Since elevated RPL19 mRNA occurred as one of a relatively small number of sequences over-expressed in prostate cancer, we hypothesized that its effect was likely to be selective rather than part of a global non-specific elevation in gene expression. Ribosomal protein L19e (RPL19) belongs to the L19E super-family of proteins and, in eukaryotes, is a component of the ribosomal large 60S subunit. The gene is expressed throughout much of evolution, particularly in eukaryotes and archaea but is absent from bacteria [19,20].
Although there is homology between sequences in rat L19 and E. coli ribosomal proteins L18, L30 and S2 [21] surprisingly, for such an apparently important gene, RPL19 has thus far received little attention. In humans, RPL19 maps on chromosome 17 at 17q11.2-q12 where it encodes 9 potential splice variants. In a series of human breast cancer biopsies, RPL19 has been reported as being expressed and co-amplified together with ERBB2 and genes PNMT, PSMB3 and NR1D1 [22]. This complex region containing multiple genes has been suggested as a possible amplicon [23,24] extending for some ~547 kb from RPL19 through STRAD3 and ERBB2 to GRB7 in the region 17q11.2-q12. Presently, no data have substantiated this speculation. In prostate cancer, amplification of erbb2 is infrequent, being reported in only 0.04% [25] to 2% [26] of cases, and therefore not a common mechanism of RPL19 over-expression. Since our initial identification of RPL19 in prostate cancer [18], its expression has been shown to define poor-prognosis colorectal cancer [27] and as a novel tumor antigen in lung adenocarcinoma [28].

Global changes in genes modulated in human prostate cancer have previously been profiled using DNA expression array analysis [29] that have detected changes in gene expression following selective up-regulation of individual target genes [30,31] or following gene-knockdown using antisense [32] or RNAi [33] technology with subsequent transformation of the malignant phenotype. The differentially-expressed genes and their associated networks have been assessed as biomarkers to segregate different prostate cancer phenotypes according to behavior and response to therapy [34]. However, an altered level of gene expression does not, ipso facto, confirm a primary role in the malignant process. Genomic instability is the hallmark of malignant transformation [35] and the effects of gain or loss of a single gene are likely to be transmitted throughout the genome with the consequence that expression of other genes becomes secondarily modulated [36]. Such changes either may have immediate and active relevance to the resulting cellular phenotype or their altered expression is passive and inconsequential. To assess the functional relevance of a particular gene, suppression of its transcription allows analysis of its immediate effects on genome-wide expression. Previously, we have transfected malignant prostatic epithelial PC-3M cells with a 436 bp-long antisense oligonucleotide to knock-down expression of FABP5 that ameliorated the malignant tumor phenotype both in-vitro and in-vivo [37]. herein, we have employed the same surgical technique of RNA interference (RNAi) with potentially greater specificity and efficiency, depending upon the particular gene being targeted [38].

Our previous data [10] indicated that expression of RPL19 might be functionally important in promoting prostate malignancy. We have now tested this hypothesis by selectively reducing RPL19 expression using RNAi. The resulting PC-3M cells exhibit an abrogated malignant phenotype both in-vitro and in-vivo when submitted to phenotypic assessment and gene expression analysis. The data support the possibility of a functional role for RPL19, acting within a spectrum of altered gene expression, in maintaining the malignant phenotype of human prostate cancer cells. Confirmation of such a scenario would allow selective therapeutic targeting of RPL19, either immunologically [28] or using small molecules, to modulate discrete subsets of cellular proteins that are key promoters of the malignant phenotype.

**Results**

**siRNA knockdown of RPL19 in parental PC-3M cells**

**Transient transfection.** qPCR analysis of the parental PC-3M cells using the primers defined in Table 1 revealed strong RPL19 mRNA expression, confirmed by nucleotide sequencing. Thereafter, transient transfection of siRNA sequences to RPL19 exon 1 (Table 2) revealed Target #1 to be the most effective sequence for RNA silencing, reducing its expression to only 7% of its initial level (Figure 1A). While the other sequences were effective, only the combination of all three simultaneously was better than Target #1, alone. Thereafter, Target #1 was used for all subsequent experiments.

**Stable transfection.** Levels of RPL19 mRNA were measured in PNT2, PC-3Mparental, PC-3Mscramble and si-RPL19-PC-3Mtarget #1 transient transfectant cells (Figure 1B). In accordance with the

**Table 1.** Primer sequences employed for qPCR identification and quantification of mRNAs.

| Primer          | Direction | Sequence                      | Amplicon Size |
|-----------------|-----------|-------------------------------|---------------|
| RPL19           | Forward   | GGGCATAGGTAACGGGAAAGG         | 149           |
| Human β-actin   | Reverse   | TCAGTACAGCCGCTGTGATACA        | 137           |
| stromelysin 1 (MMP3) | Forward   | AAATCCCTCAGGAAACGCTGA         | 139           |
| stromelysin 2 (MMP10) | Forward   | CAGGACACAGTGGTTGCTCAT         | 101           |
| collagenase 3 (MMP13) | Forward   | GGTGGTATGTAATCTCGTGAGG        | 139           |
| Fas (TRIF6) associated factor 1 (FAF1) | Forward   | CTTCAGGCTTCCGACTGTA           | 225           |
| nuclear factor of kappa light polypeptide gene enhancer in B-cells inhibitor, alpha (NfκBIA) | Forward   | TCCGAGACCTCGAGAAAT            | 143           |
| STON2           | Forward   | AGCACTGGTGATGTTAA             | 90            |
|                 | Reverse   | GGTCAATGTGAGGCCGCTCT          |               |

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previous study [33], expression in PC-3M
scramble cells was set at unity and relative expressions in the other cell-lines were compared as fold-differences. RPL19 expression in PC-3M was 4.9 times greater than that of the PNT2 cells and consistent with our previous studies confirmed by Northern blot analysis [18]. In the si-RPL19-PC-3Mclone ST-3 transfectant cells, expression of RPL19 was reduced to only 1.3 times greater than the PNT2 cells. PC-3Mscramble cells revealed a 2.3 fold reduction in RPL19 when compared to PC-3Mparental, although this value was not statistically significant. Single cell cloning [33] followed by qPCR and Western blotting confirmed si-RPL19-PC-3Mclone ST-3 expressed the lowest levels of RPL19 mRNA and protein. This clone of cells was thereafter employed for detailed phenotypic analysis.

Growth characteristics of si-RPL19 cells in-vitro

Clones of transfected si-RPL19-PC-3M cells grown under standard conditions exhibited differences in morphology (Figure 1C). Compared to PC-3Mparental cells, si-RPL19-PC-3M cells were generally less adherent to substrate. However, these cells maintained an ability to proliferate and could be successfully subcultured, although a large proportion of the cells remained in suspension. Other si-RPL19-PC-3M cells showed an increase in multinucleate forms, suggesting impaired completion of mitosis. Proliferation assays (Figure 2A) revealed that during the logarithmic phase of growth, the rate of cell division by the si-RPL19-PC-3Mclone ST-3 transfectant cells was not significantly affected ($p<0.05$) when compared to PC-3Mparental and si-PC-3Mscramble. The ability of si-RPL19-PC-3Mclone ST-3 cells to invade an extracellular collagenous matrix (ECM) was compared to that of the PNT2, PC-3Mparental and PC-3Mscramble cell-lines (Figure 2B). The number of cells that invaded through the ECM were: (PNT2) 0.6 ± 0.6, (PC-3Mparental) 279 ± 33.7 and (PC-3Mscramble) 317 ± 28.3 ($p<0.001$). The si-RPL19-PC-3M cells exhibited a comparatively poor invasive potential at only 60 ± 10.7 transmigrating cells ($p<0.001$). Thus, silencing RPL19 reduced the invasive potential of PC-3M cells approximately 5-fold. Endogenous (basal) levels of apoptosis within the PC-3Mparental and PC-3Mscramble cells (Table 3 and Figure 2C) were similar to those obtained during comparable studies of the PKRζ gene [33]. Basal levels of apoptosis in the four cell-lines were not statistically different ($p>0.05$). Although sensitivities of the PC-3Mparental and PC-3Mscramble cell-lines revealed no statistically significant differences ($p>0.05$), indicating continued low-level expression of RPL19 in the majority of tumor cells. Detection of small amounts of RPL19 protein in some tumor cells is considered to represent clonal variation resulting from continued low-level expression of the gene, rather than its total inhibition, as identified by qPCR of the cells in vivo and the results of the Western blotting studies. While expression of mRNA and corresponding protein in prostatic epithelium are not always concordant [39], apparent discrepancies between in-vitro and in-vivo studies may be due to the in-vivo effects of a surrounding stromal matrix affecting tumor cell adhesion or to other influences including growth factors modulating individual low-level gene expression [40–42].

Tumorigenicity and RPL19 protein expression in-vivo

In all groups of animals, tumors became apparent on day 2 following inoculation (Table 4). However, more appeared sooner in the PC-3Mparental (3/8) and PC-3Mscramble (4/8) groups. In the two transfectant clone groups, tumors took longer to appear (2/8 tumors in animals carrying the si-RPL19-PC-3Mclone ST-3 cells and 1/8 tumors in animals carrying the si-RPL19-PC-3Mclone ST-2 cells). Initially, all tumors were similar in size. After 7 days the PC-3Mparental and PC-3Mscramble groups developed larger tumors than two transfectant groups (Figure 2D). At autopsy, 15 days after inoculation, a significant difference ($p<0.001$) was apparent in the mean weights of the control and RPL19 knockdown tumors (Figure 2E). PC-3Mparental exhibited a wide range in tumor weight, one animal producing a tumor of 810 mg in 15 days, the maximum allowed by the Project License. Conversely, another animal developed a tumor of only 10 mg. A similar phenomenon occurred within the PC-3Mscramble group with tumors ranging from 10-140 mg. The final weights of the PC-3Mparental tumors were not significantly different from those of the PC-3Mscramble group (Mann-Whitney U Test, $p>0.05$). Thus, si-RNA suppression of RPL19 affected the size of the tumors generated in-vivo ($p<0.05$) but not on their latency. No micrometastases were identified at autopsy or on subsequent histopathological examination of the excised tissues.

Immunohistochemistry of tumor xenografts detected strong expression of RPL19 protein in both the PC-3Mparental and si-PC-3Mscramble cells (Figure 2F). Knockdown cell lines si-RPL19-PC-3Mclone ST-1 and si-RPL19-PC-3Mclone ST-3 exhibited comparatively little staining, indicating continued suppression of the RPL19 gene in the majority of tumor cells. Detection of small amounts of RPL19 protein in some tumor cells is considered to represent clonal variation resulting from continued low-level expression of the gene, rather than its total inhibition, as identified by qPCR of the cells in vivo and the results of the Western blotting studies. While expression of mRNA and corresponding protein in prostatic epithelium are not always concordant [39], apparent discrepancies between in-vitro and in-vivo studies may be due to the in-vivo effects of a surrounding stromal matrix affecting tumor cell adhesion or to other influences including growth factors modulating individual low-level gene expression [40–42].

Comparative gene expression profiling of si-RPL19-PC-3Mclone ST-3 cells

Genome-wide expression profiles obtained from DNA oligonucleotide microarrays (unmodified Agilent Human Genome 44K) were employed to identify genes modulated following RPL19 knockdown. Comparison of genes expressed by PC-3Mparental and PC-3Mscramble cell-lines revealed no statistically significant differences ($p>0.05$), indicating that the transfection technique was not responsible for appreciable off-target effects that might bias the experimental data. A total of 916 DNA sequences, representing 768 genes, were identified as differentially expressed ($p<0.05$, Benjamini and Hochberg multiple testing correction applied). Of these, 404 were enhanced and 364 down-regulated. Within that data set, 184 different genes were modulated at least four-fold, 62 being up-regulated and 122 down-regulated. The top 50 differentially-expressed genes in these two categories are summarized in Supporting Information Tables S1 and S2 and graphically

| Target | Sequence | Position in gene sequence | Exon silenced | Variants affected |
|--------|----------|--------------------------|---------------|------------------|
| #1     | AGGCTATCAAAGATGGCTG | 15 | 11 | a, b, c, d, e, g |
| #2     | AAAAACGGGATTCTCATGGA | 43 | 18 | a, c, d, f, h |
| #3     | AGATACCGTGAACTAAGAA | 86 | 15 | a, b, c, d, e |

This Table identifies the particular exon silenced and the alternative splice variants predicted to be affected.

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Expression data derived from the arrays were validated by qPCR providing independent quantifiable evidence of the magnitude and direction of change of individual genes. The observation that only 768 genes were modulated following RPL19 knockdown, with the levels of mRNA for a wide range of proteins either maintained or elevated, suggests that ribosomal protein RPL19 is differentially involved in protein synthesis rather than affecting all cellular protein synthesis in a non-specific manner.
Figure 2. Growth characteristics of si-PC-3Mclone ST-3 cells in-vitro and in-vivo. A. Relative growth of cell-lines in monolayer culture revealing no statistical difference in the rate of proliferation between the knockdown cells (si-RPL19-PC-3Mclone ST-3) and that of PC-3Mparental cells. B. Invasion assay in-vitro comparing the same populations of cells as shown in (A) and revealing an 83% decrease in the invasive activity of the RPL19 knockdown cells relative to PC-3Mscramble cells. C. Resting levels of apoptotic indices were not significantly different in the benign (PNT2), parental (PC-3M) or knockdown cells. After challenge by camptothecin, no change was identified in the PC-3Mparental or si-RPL19-PC-3Mclone ST-3 cells. While an increase in apoptosis was found in the benign (PNT2) and scramble-transfected cells, these were not significant. D. Growth of tumor cells in vivo by estimated volume revealed a highly significant (p<0.005) suppression of growth by two of the stable transfectant clones, when compared to the PC-3Mparental and PC-3Mscramble growths. Growth of PNT2 cells is not included since we have already shown [33] the growth of tumors to be infrequent, particularly over the time-span of these experiments. E. Analysis of tumor weights comparing the same populations of cells as those shown in (A) and revealing an 83% decline in the invasive capacity of the si-RPL19-PC-3Mclone ST-3 cells (iii) expressed RPL19 heterogeneously and at only very low levels. (Magnification ×350).

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Functional enrichment analysis identifying some 20 Gene ontology (GO) biological process terms and three molecular function terms (Supporting Information Table S3) to be significantly associated (p<0.001) with the knockdown (p<0.001). Additionally 13 KEGG pathways had a high level of expression differentially between RPL19-PC-3Mclone ST-3 and PC-3Mscramble (Supporting Information Table S4). Ingenuity pathway analysis was used to identify significant biological networks and pathways in which the genes expressed differentially as a consequence of RPRKC-ζ knockdown were involved. The top five ranked interlinked pathways (Supporting Information Table S5) and the three Gene ontology (GO) molecular function terms (Supporting Information Table S6) are highly significant (p<10^-7) with respect to genes differentially expressed after RPL19 knockdown.

Ribosomal protein genes. The hypothesis that siRNA-induced down-regulation of RPL19 might be compensated by modulation of other ribosomal proteins was addressed by assessment of the relative expression of the mitochondrial large ribosomal protein gene sequences (n=71) and the cytoplasmic large ribosomal protein gene sequences (n=136) to discover whether up-regulation of a gene already expressed or neoexpression of a previously silent ribosomal protein gene had occurred. Of the latter cohort, 44 genes identified known RPs, 7 were RP-like and 5 were RP pseudogenes. The number of sequences representing each gene ranged from one (19 genes) to 10 (RPL21). RPL19 was identified by a single sequence. According to SCOP (Structural Classification of Proteins, latest release 9th November 2010, http://scop.mrc-lmb.cam.ac.uk/scop) RPL19 is a member of the protein superfamily of translation proteins containing the SH3-like barrel structural domain within the Class comprising all beta proteins. The family also contains ribosomal protein genes RPL14c, RPL21c and RPL24p and the G-terminal domain of RPL2 (http://supfam.org/SUPERFAMILY/cgi-bin/scop.cgi?unid=30104). Alternatively, RPL19 protein could be replaced by RPL29 or RPL39e, being structurally similar members of the ζ helical group of globular RPs with extended tails able to bind mRNA [43]. Although fluctuations occurred in the levels of expression of individual RPL gene sequences following RPL19 knockdown, these were not significant, including that of ribosomal protein gene RPL23A also located on chromosome 17q11.2. Only expression of mitochondrial MRPL42 was significantly down-regulated (p<0.05). No enhanced expression of any RP gene was detected. Thus, inhibition of RPL19 with loss of RPL19 protein was not compensated by a different RP gene. Conversely, the effects of reducing RPL19 could be mediated by the coding-independent function of the gene or its pseudogene mRNAs [44].

Glycosyltransferase genes. Transformation of epithelial cells from a benign to a malignant phenotype is often accompanied by structural changes in the oligosaccharide domains of cellular glycoproteins and glycolipids [45]. Particularly, expression of sialylated and β-1,6 branched N-linked oligosaccharides are required for cancer cell invasion and metastasis [46]. The key enzyme in this process is mannosyl (α-1,2)-glycoprotein β-1,6-N-acetylglucosaminyltransferase encoded by gene MGAT3 and regulated by signaling pathway RAS-RAF-MAPK. Together with PTEN, MGAT3 regulates the membrane dynamics of PI3K/Akt signaling to promote the invasive malignant phenotype [47]. In the event that malignancy is reduced following manipulation of cellular phenotype, changes in cell-surface oligosaccharide structures are postulated to occur. Such changes, mediated by glycosyltransferases may be evidenced by altered expression of the corresponding genes. Of the 768 genes differentially-expressed, only two glycosyltransferase genes were significantly affected following RPL19 knockdown (Supporting Information Table S7). Unlike the spectrum of glycosyltransferases modulated following si-RNA knockdown of PRKC-ζ in PC-3M cells [33], no change was apparent in sialyl- or fucosyl-transferase genes. However, a 4-fold reduction was identified in the level of MGAT4 (p<0.05) that encodes the enzyme mannosyl (α-1,3,4)-glycoprotein β-1,4-N-acetylglucosaminyltransferase and is involved in mediating glycosylation of the proteins encoded by SLC43A3 (proteoglycan 2), SLC14A1 (urea transporter) and SLC38A1 (sodium/calcium exchanger), thereby controlling their cell-surface expression. Indeed, all three latter genes were modulated following RPL19 knockdown. Conversely, a 2–3-fold increase was identified in the level of GALNACT-2 (p<0.05) that encodes the enzyme chondroitin sulfate N-acetylgalactosaminyltransferase 2 and transfers N-acetylgalactosamine (GalNAc) from UDP-GalNAc [48] to chondroitin, chondroitin sulfate, preferentially to complex oligosaccharides containing β1→4 linkages[49], such as those generated by MGAT4.

Ion channels and associated genes. The malignant phenotype of prostatic epithelial cells can be modulated by differential expression of ion channels [50–52]. Studies from this laboratory [52] and elsewhere [53] have established a functional relationship between voltage-gated ion channels and the invasive

| Cell-line | Basal level of apoptosis (%) | Level of apoptosis following camptothecin (%) | Student’s t test (p) |
|-----------|-----------------------------|--------------------------------------------|-------------------|
| PNT2      | 5.68±3.4                    | 24.5±1.4                                   | ≤0.0001           |
| PC-3Mparental | 5.3±1.4                  | 4.7±1.4                                    | >0.05             |
| PC-3Mscramble | 7.6±1.9                  | 21.5±1.4                                   | <0.001            |
| si-RPL19-PC-3Mclone ST-3 | 3.9±0.9               | 4.43±1.4                                   | >0.05             |

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Table 4. Incidence and latency period of tumors produced by transfectants in nude mice.

| Cell-line                        | No of animals inoculated | Incidence of tumors* | Median latent period in days (range) | Mean weight of tumors (mg/2) ** |
|----------------------------------|--------------------------|----------------------|-------------------------------------|-------------------------------|
| PC-3Mparental                    | 8                        | 8                    | 100                                 | 5.25 (2-12)                   | 337.5±266.5                  |
| PC-3Mscramble                    | 8                        | 8                    | 100                                 | 4.275 (2-12)                  | 120±83.7                     |
| si-RPL19-PC-3Mclone ST-3         | 8                        | 8                    | 100                                 | 6.875 (2-12)                  | 36.0±35.0                    |
| si-RPL19-PC-3Mclone ST-3         | 4                        | 4                    | 50                                  | 4.25 (2-5)                    | 7.5±13.9                     |

*Tumor incidence is the percentage of mice with tumors/total number of inoculated animals.

**The final weights of the si-RPL19-PC-3Mclone tumors were significantly less than the PC-3Mparental and the PC-3Mscramble tumors (Mann-Whitney U Test, p<0.05).

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Phenotypic gene expression in prostatic malignancy

Protein expression. The finding that the rate of cell proliferation did not decline following gene knockdown suggested that global suppression of protein synthesis was unlikely to have occurred despite expression of an individual ribosomal protein being significantly reduced. However, a differential effect was identified with respect to individual proteins (Figure 5), exemplified by AGR2 that was abrogated in the si-RPL19-PC-3Mclone ST-3 cells while expression of ERBB2 was simultaneously enhanced. The observation that individual proteins were differentially affected suggests the biological effects of reducing RPL19 to be gene-specific and protein-specific rather than a global down-regulation of protein synthesis. The enhanced level of ERBB2 provides additional evidence against a common amplion in chromosomal region 17q11.2-q12 since expression of RPL19 and ERBB2 were divergent (Figure 5).

Hsp-27 expression and phosphorylation status. Western blotting confirmed the level of total Hsp-27 protein to be lower in the si-RPL19-PC-3Mclone ST-3 and si-EAB55-PC-3M cells than in the PC-3Mparental PC-3Mscramble and the si-PRKC-ζ-PC-3Mclone cell-lines (Figure 5). A global reduction in site-specific phosphorylation of si-RPL19-PC-3Mclone ST-3 cells was also identified when compared to PC-3Mparental and PC-3Mscramble in contrast to the effect of knocking-down PRKC-ζ [33].

Glycoconjugate expression. Lectin histochemistry, employed to test the hypothesis that RPL19 knockdown would modulate the profile of sialylated glycoconjugates, revealed no qualitative difference in expression of Neu5Acα2→3Gal- and Neu5Acα2→6Gal- (using Sambucus nigra and Maackia amurensis, respectively) when the PC-3Mparental and si-RPL19-PC-3Mclone ST-3 cells were compared. Staining was abolished in all cell-lines following neuraminidase digestion, confirming specificity of sialic acid expression. Similarly, the lectins from Ulex europaeus, Lotus tetragonolobus and Aleuria aurantia revealed no changes in terminal fucosyl linkages. Thus, suppression of the metastatic phenotype by knockdown of RPL19 did not involve appreciable loss of sialic acid from the cell surface.

Discussion

This study provides evidence that posttranscriptional silencing of RPL19 using RNAi not only abrogates the malignant phenotype of PC-3M prostate cancer cells but is selective with respect to transcription and translation of other genes. In prostate cancer, expression of RPL19 is significantly elevated, functionally involved in maintaining the malignant phenotype and hence a potential target for therapeutic intervention. Despite its involvement in ribosome structure and function, the data show that the effects of reducing RPL19 are not global but restricted to a defined cohort of genes and proteins. This observation supports the accumulating evidence of eukaryotic ribosomal specialization in which loss of
Figure 3. Graphical representation of gene expression modulated following \textit{RPL19} knockdown. Heat map of top 50 genes up-regulated and top 50 genes down-regulated following expression-profiling of mRNA expressed by \textit{si-RPL19-PC-3M} clone \#3 cells when compared to \textit{PC-3M} parental cells using \textit{PC-3M} scramble cells as the common denominator. Hierarchical clustering is shown. Green indicates genes over-expressed in a sample compared to scramble-transfected cells. Red indicates genes down-regulated in the sample when compared to scramble-transfected cells. Corresponding numerical data are presented in Supporting Information Tables S1 & S2.

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function phenotypes of individual ribosomal proteins is associated with changes to specific signaling pathways and tissues [57] suggesting that core ribosomal proteins may contribute differentially to translation of distinct subpopulations of mRNAs [58]. Within ribosomes, the role of RPL19 remains undefined. Nevertheless, importance of the gene may be inferred from the number of its paralogous sequences maintained within the eukaryotic genome [59] and the finding that expression of RPL19 is one of the most stable and consistent genes within the human genome [60,61].

Analysis of the 768 genes modulated following RPL19 knockdown revealed a genetic profile distinct from that obtained after siRNA reduction of PRKCZ [33] or FABP5 [62] in the same PC-3M cells. The cohort of modulated genes did not contain any cell cycle-associated genes [54], DNA-binding genes (e.g. RAD51) or transcriptional activation genes (e.g. Id-1) we have already reported in aggressive primary prostate cancers [63,64]. Furthermore, the affected gene-networks did not involve cell adhesion genes or other ribosomal protein genes identified in metastatic breast cancer [65]. However, AGR2 was down-regulated (>10-fold, \( p<0.001 \)) in the knockdown cells, consistent with our previous finding in non-malignant prostatic epithelium [55]. Although the levels of some ion channel and glycosyl transferase genes were appreciably modulated, individual members of these cohorts were different from those identified following RNAi gene-knockdown of PRKCZ [33] providing additional evidence that gene expression is heterogeneous within the benign phenotype.

While enhanced expression of some RP genes has been reported in other human malignancies [66], including lung [28], colorectal [67], prostate [68] and RPL19 in breast cancer [69] this is the first report to define a functional role for RPL19 in the malignant phenotype. Although RPL19 protein is an integral component of the large 60S subunit of eukaryotic ribosomes [59,70] its ribosomal function has not been defined. Nevertheless, reduction in its expression sufficient to modulate the behavioral phenotype from malignant to benign did not involve a detectable alteration in cell proliferation or apoptosis indicating that the phenotypic effects were not simply due to the target cells becoming compromised either metabolically or by diminished protein synthesis. In the absence of non-specific global effects, the data indicate that reducing RPL19 expression affects discrete populations of genes and proteins, thus shifting the balance of gene expression from a malignant to a benign phenotype.

Intuitively, loss of RPL19 protein might be expected to cause a general decline in ribosome biosynthesis with compromised functionality and commensurate loss in protein synthesis. Under such circumstances, cell proliferation would have decreased without specific effects on particular cellular functions. In contrast,
the data indicate the effects to be selective with respect to cell adhesion, stromal invasion and tumorigenesis. Although the role of individual RPs in determining the cellular phenotype of eukaryotic cells remains unclear, current evidence reveals mutations within individual ribosomal proteins to be associated with specific changes in cellular phenotype [71,72] rather than a general down-regulation of protein synthetic activity. Examples emerging within other fields of protein biology indicate that alternative genes may be recruited to replace defective or deficient proteins [73–75]. Although such mechanisms would be important to maintain structure-function relationships within complex organelles, no such examples have been reported to compensate for deficient ribosomal proteins. Subsequent to the loss or replacement of an individual ribosomal protein, the functional activity of the modified organelles would not be identical to that of the original structure, thus providing a drive towards adaptation and evolution of a novel phenotype [76,77]. In addition to protein biosynthesis, many RPs also fulfill extra-ribosomal functions, particularly regulating the quality of gene expression through coupling transcription mechanisms with the processing and transportation of mRNAs [78,79]. Such effects are stochastic and cannot be predicted because of the complexity gene interactions [80]. Nevertheless, mathematical models are emerging to analyze the effects of insertions and deletions in protein-protein interaction networks and the global changes consequentially induced in cellular structure and function [31–33]. Despite protein synthesis being a general function of ribosomes, the precise function of each ribosome depends upon its complement of ribosomal proteins, ribosomal RNAs (rRNAs) and a range of ribosome-associated proteins (PARs) [84]. Ribosome biogenesis is complex and highly regulated [85,86]. Continuity of the cell-cycle depends upon fidelity of ribosome biogenesis and ceases if ribosome biogenesis becomes impaired [10,37–39], leading to a variety of ribosomopathies [90]. Such data provide evidence that structurally-defective ribosomal components (rRNAs, RPs or PARs) cause disruption of a cell’s translational apparatus [91], resulting in alterations to cellular phenotype [92] with the consequence that small changes in molecular structure may cause significant alterations in ribosomal function.

Herein we confirm a functional role for RPL19 in promoting the malignant phenotype of human prostate cancer cells. Despite significant reduction in the levels of RPL19 mRNA and protein, the finding that cell proliferation was not demonstrably affected challenges the supposition that RPL19 protein is essential for ribosomal structure and/or function and suggests a level of adaptation within ribosomal protein function that enables global protein synthesis to be maintained despite loss of a core ribosomal component. If RPL19 is not a critical component of ribosome structure and/or function, its importance to the malignant phenotype may be related to its extra-ribosomal activities, with the implication that there is no necessity for protein substitution or adaptation at the ribosomal level. Our finding that the patterns of genes and their associated networks modulated by RPL19 knockdown are distinct from the patterns following PRKCZ knockdown in identical cells is consistent with two propositions: First, that loss of individual ribosomal proteins is associated with specific alterations in cellular phenotype. Second, that the non-malignant phenotype is not defined by a single immutable pattern of gene expression but is in flux [93] in the same manner that the patterns of genes expressed in malignant cells are heterogeneous [94,95]. The possibility of flux between metastable gene networks raises the exciting possibility of therapeutically stabilizing a benign phenotype generated by modulating expression of a key gene and hence constraining a malignant phenotype while leaving non-malignant genomes unaffected.

Materials and Methods

Cell lines

Human prostate cell-lines PNT2 (benign) and PC-3Mparental (highly malignant) are identical to those described previously [33]. PNT2 cells are non-malignant, androgen-independent and derived from SV-40 immortalization of normal prostatic epithelial cells [96]. PC-3M cells are malignant, also androgen-independent and derived from the bone marrow metastasis of a 62 year-old man [97]. These cells exhibit a high incidence of tumorigenesis and metastasis when xenografted into nude mice [98]. Both cell-lines are histogenically the closest currently available having contrasting behavioral phenotypes and hence the most appropriate as comparators. Gene knockdown derivatives of the PC-3M cell-line si-FABPS5-PC-3Mclone 3 [62] and si-PRKCZ-PC-3Mclone 3 [33] described in comparable studies and were employed to reveal similarities and differences in gene-expression following abrogation of the malignant phenotype in PC-3M cells using an identical technique. All cell-lines were grown as monolayer cultures in RPMI 1640 (Invitrogen, Paisley, UK) supplemented with 10% (v/v) fetal calf serum (FCS, Invitrogen), penicillin (100units/ml), streptomycin (100 µg/ml), and L-glutamine (2 mM). Media for the culture of all subsequent transfected cell-lines were also supplemented with 1 µg/µl Geneticin (Sigma).

siRNA Knockdown of RPL19 in PC-3M cells

Transient transfection. Transient transfections were performed by the reverse transfection technique using siPORT NeoFX Transfection Agent (Ambion, Warrington, UK). Three transfections were initially assessed as potential targets for stable transfection to silence variant “c”, the NM version of the RPL19 gene (NM_000981). All three sequences were potentially capable of silencing seven of the alternative eight splice variants (a, b, c, d, e, f, g, h) of RPL19. The alternative 168 bp variant “i” was incomplete since it did not contain the target. The transfection targets, listed in Table 2, were BLAST-searched and showed
homology and similarity only to RPL19. Sequences to targets were designed using Ambion’s online target design algorithm and purchased from Ambion was also included, pre-annealed at a concentration of 20 nmol.

A negative control siRNA was also included that comprised a nucleotide sequence similar in composition to that of the siRNA but not homologous to any known gene of interest and purchased pre-designed from Ambion. This “scramble” sequence was used to discount non-specific changes in gene expression profiles due to siRNA delivery. Preliminary experiments optimized transfection conditions for PC-3M cell lines. Reverse transfection was performed in a 96 well plate format. Cells were seeded at a density of 8x10⁵ cells/well. The short strand RNA (sRNA) oligonucleotide sequences were then diluted in a reduced serum medium (OPTI-MEM 1; Invitrogen, Paisley, UK) to a final concentration of 30 nM. This was then overlaid onto the cells that were then incubated at 37°C in 100% humidity in 5% CO₂/air for 24 hours. Transfection of the RNA oligonucleotide sequences into the cells occurred spontaneously as the cells adhered to their substrata.

Stable transfection. After transient transfection had identified Target #1 as the most successful to silence RPL19 expression, this sequence was used to generate a hairpin siRNA. The following oligonucleotides were purchased from Ambion:

**Top Strand:**
5'-GATCCGCTCATCAAGAGTGCGCTTGTCAGAGAA-3'

**Bottom Strand:**
5'-AGCCTTAAGCTATCAAGAGTGGGTCGCCTTCG-3'

The default Ambion loop sequence, TTCAAGAGA, was used to complete the hairpin structure. The siRNA expression vector kit used was pSilencer® 4.1-CMV neo (Ambion). Top and bottom strands of the siRNA hairpin oligonucleotide were diluted to 1 µg/µl in TE buffer and annealed in 50 µl solution according to the manufacturer's instructions. The annealed siRNA template was ligated into the pSilencer® 4.1-CMV vector using T4 DNA ligase (5 U/µl) and the products cloned into DH5α cells (Invitrogen).

Transformed cells were grown for 16 hours on LB plates containing 100 µg/ml ampicillin at 37°C. A negative control of non-transformed competent cells was also included. Clones were picked and the DNA plasmid isolated using a Qiaprep spin Miniprep Kit (Qiagen, Crawley, UK). Isolated plasmids were digested with BamHI and HindIII (New England Biolabs, Hitchin, UK) and the presence of the siRNA 55 bp insert was confirmed by sequencing prior to the siRNA expression vector being used to transfect recipient prostate cancer cell lines. Orientation of the insert was confirmed by DNA sequencing (Lark Technologies, Essex, UK) using internal sequence primers.

**Transfection of siRNA RPL19 silencing construct and control.** 1.5x10⁵ PC-3M_parental cells were transfected with pSilencer® 4.1 CMV RPL19 siRNA (1 µg) using SiPORT XP-1 (3 µl) reagent (Ambion, Warrington, UK) in 6-well-plates (35 mm diameter). 24 hours after transfection, 500 ng/ml of G418 was added to medium RPMI1640 for selection. After 9-10 days selection, individual colonies from single cells containing stable clones were isolated using ring cloning and transferred into 24-well plates with medium containing G418 at 500 ng/ml. Simultaneously, 1.5x10⁵ PC-3M_parental cells were transfected with pSilencer® 4.1 CMV-scramble-insert (1 µg). Thereafter, these cells were cultured, cloned and employed as the controls to assess changes in expression of genes and proteins by the knockdown cells.

**RNA extraction and cDNA synthesis.** Total RNA was extracted with RNeasy Mini Kits (Qiagen). Total RNA concentration was measured using a NanoDrop instrument (Labtech, Ringmer, UK) and RNA integrity assessed with a 2100 Bioanalyzer (Agilent, Santa Clara, USA). The RNA integrity number (RIN) for all RNA used exceeded 9.0. First strand cDNA was synthesized from 0.5 µg total RNA using AffinityScript™ Multiple Temperature cDNA synthesis kits (Stratagene, La Jolla, USA) according to the manufacturer’s protocol.

**Quantitative Real-Time PCR (qPCR).** RPL19 mRNA expression levels were quantified by qPCR and normalized relative to human β-actin mRNA expression. An MX3005P Real Time PCR machine (Stratagene) was used for all reactions. Reaction volumes were in 25 µl comprising 12.5 µl Stratagene’s Brilliant® SYBR® Green Master Mix (2X), 0.5 µM of both forward and reverse primers and 1 µl cDNA and 11.5 µl water. Primers for qPCR were designed to span exon/exon boundaries within the mRNA to avoid amplification of genomic DNA. Primers designed for RPL19 and human β-actin are listed in Table 1. Both primers were optimised at 60°C. Cycling conditions for the reaction were: 95°C for 15 minutes, then 40 cycles at 94°C for 15 seconds, 63°C for 30 seconds, plate read and 72°C for 30 seconds with a final extension at 72°C for 10 minutes. Melting curves were generated to detect primer-dimer formation and to confirm gene-specific peaks for targets.

**Growth characteristics and invasiveness of si-RPL19 cells in-vitro.** An assay was established to identify the effect of RPL19 suppression on cellular proliferation. The relative growth rates of PNT2, PC-3M_parental, PC-3M_scramble and si-RPL19-PC-3M clone ST-3 transfectant cells were measured by proliferation assay. Exponentially-growing cells were seeded in triplicate sets at a density of 1x10⁵ cells/ml/well in 24-well plates. Over 10 days at 24–48 hour intervals, cell proliferation was calculated by measuring the increase in cell numbers in each replicate using a conventional MTT assay [99]. Apoptosis was quantified using flow cytometry. Cells from PNT2, PC-3M_parental, PC-3M_scramble and two si-RPL19-PC-3M clones were seeded in which 2x10⁵ cells/ml in 75 cm² tissue culture flasks and the assay started prior to cells reaching confluence. Duplicate flasks were established in which cells were exposed to 4 µM camptothecin (Sigma-Aldrich) dissolved in DMSO for 24 hours before harvesting. Camptothecin, a potent inhibitor of topoisomerase I, induces apoptosis in a dose-dependent manner in-vitro [100,101]. Cells were harvested by trypsinization, washed twice with PBS and re-suspended in buffer from the BioVision Annexin V-FITC kit in a 5 ml flow cytometry tube. AnnexinV-FITC (5 µl) and propidium iodide (10 ng in 5 µl aqueous solution) were added and the tubes incubated for 10 minutes in darkness at 4°C. Quantitative analyses of apoptotic cell levels were performed using an Epics Flow Cytometer (Beckman Coulter). The procedure was performed three times using biological replicates. Invasiveness of the si-RPL19 transfectants was assessed in-vitro [32]. At 24-hour intervals, following fixation and staining with Crystal Violet (Sigma-Aldrich, St Louis, USA), invasion was measured by counting the number of cells transmigrating the membrane to its under-surface [33].

**Tumorigenicity and RPL19 protein expression in-vivo.** All studies were performed under the conditions of UK Home Office Project License PPL 40/2270 [33]. Tumorigenicity was assessed by injecting cells (2x10⁶ cells in 0.2 ml PBS) into a single subcutaneous site in the right shoulder of 8 week old male Nu/nu mice (Harlan Ltd., Oxon, UK). Four groups of cells were assessed: PC-3M_control, PC-3M_scramble and si-RPL19-PC-3M clones #1 and #2. Of these two, si-RPL19-PC-3M clone ST-3 exhibited the most pronounced suppression of RPL19 and was used in the microarray and invasion assay experiments.

**Table 1. Both primers were optimised at 60°C. Cycling conditions for the reaction were:**

- 95°C for 15 minutes, then 40 cycles at 94°C for 15 seconds, 63°C for 30 seconds, plate read and 72°C for 30 seconds with a final extension at 72°C for 10 minutes.
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si-RPL19-PC-3Mg2 exhibited suppression mid-way between that of si-RL19-PC-3Mg1 and si-RL19-PC-3M scramble. Tumor growth was monitored twice-weekly by measuring the largest (a) and smallest (b) superficial diameters. Tumor volume (V) was then calculated using the formula \( V = a \times b^2/2 \) [102]. When any tumor reached the maximum size allowed under the conditions of the Home Office Project Licence PPL 40/2270, all mice were sacrificed. Each animal was submitted to autopsies to identify appearance of metastatic tumor nodules. Subcutaneous primary tumors together with heart, liver and lungs were removed and weighed. All tissues were processed and embedded in paraffin wax. Histological sections cut at 4 μm and stained with Gill’s hematoxylin for microscopic examination.

Expression of RPL19 protein in human prostate epithelial cells grown as xenografts in nude mice was detected using a mouse monoclonal antibody (Abnova, Taiwan; #H0000 6143-MO1) diluted to 1:1000 in REAL antibody diluent (Dako, cat. no. S2022). Prior to staining, antigen retrieval employed PT-Link with EnVision FLEX, high pH target retrieval solution. Staining was performed on a Dako Autostainer using a labeled polymer-HRP detection system (Dako, EnVision FLEX, K8000). Immunostained sections were counterstained with hematoxylin, dehydrated and mounted. Negative controls comprised duplicate tissue sections processed identically but with replacement of the primary antibody by a 1% (w/v) solution of bovine serum albumin. Specimens were considered positive only when at least 5% of the epithelial cells (either normal or malignant) unequivocally expressed RPL19 staining [103]. This cut-off was the same as that used to distinguish positive and negative immunohistochemical staining in our previous studies [104,105]. Staining was assessed as negative, weakly positive or only focally positive (low-level expression), or strongly positive (high-level expression) and scored as 0, 1, 2 or 3, respectively.

Microarray analysis

Microarray validation. Gene expression profiles were validated in the knockdown cells using qPCR to confirm the expression level of NFKBIA, TNFSF6, MMP2 and MMP10 (Supporting Information Figure S1) in addition to PLAT, HSPB1, CDKN2C and FOXA2, previously employed to validate these arrays [33] when normalized against human β-actin. The primers and amplicon sizes are listed in Table 1. All annealing temperatures were 60°C and cycling conditions as described previously [33].

Gene microarray and expression analysis. The effect of suppressing RPL19 by gene knockdown on whole genome expression profiles was investigated using two-color Agilent Human genome 44k microarrays. Each hybridization was a distinct biological replicate. The design incorporated five cell-lines treated as fixed biological factors: PNT-2, PC-3M parental, si-RPL19-PC-3Mg2, si-RPL19-PC-3Mg1 and PC-3M scramble. In addition to PLAT, HSPB1, CDKN2C and FOXA2, previously employed to validate these arrays [33] when normalized against human β-actin. The primers and amplicon sizes are listed in Table 1. All annealing temperatures were 60°C and cycling conditions as described previously [33].

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Phenotypic gene expression in prostatic malignancy

Protein expression. To analyse the effect of reducing RPL19 expression on the protein-synthetic function of ribosomes, Western blotting was performed using a range of commercially-available antibodies (Supporting Information Table S8). Proteins were extracted from ~1×10^7 cells from each line. Cell pellets were suspended in 1 ml of CelLytic-M lysis buffer (Sigma C2978) containing 10 μl protease inhibitor cocktail (Sigma P8340), 10 μl PMSF (0.1 mg/ml), Na₃VO₄ (1 mM) and NaF (1 mM). Protein concentrations were determined by Bradford assay (BioRad kit 500-0006). Aliquots containing ~10 μg cell lysate proteins were separated electrophoretically at 150 V in 12.5% (w/v) polyacrylamide NextGel quick-cast separating gels (Amresco, Solon, OH). Separated proteins were transferred onto PVDF membranes (GE Healthcare, RPN303F), at 100 V for 1 hour, blocked with a suspension of powdered dried milk in PBS (100 mM, pH 7.6) before incubation at 4°C with primary antibodies. After washing and incubation with the corresponding anti-(mouse Ig)- or anti-(rabbit Ig)-HRP antiserum at 1:10,000 dilution for 1 hour, washing and incubation in ECL Plus reagent (GE Healthcare, RPN 2133) for 5 minutes, exposure to Amersham Hyperfilm (GE Healthcare, 28906839) for 5 seconds before being developed and fixed. To quantify protein expression, membranes were incubated with an anti-beta actin mouse monoclonal antibody for 30 minutes. Bound anti-actin antibody was detected as described. A strong single band at 42 kDa was observed in all cases.

Hsp-27 expression and phosphorylation status. Hsp-27 is an independent biomarker of the aggressive malignant phenotype of human prostate cancer [105]. Although no...
functional relationship has been reported between Hsp-27 and PKC-ζ, it was hypothesized that amelioration of malignancy following RPL19 knockdown would be accompanied by a reduction in the level of Hsp-27 expression. Expression of Hsp-27 is a validated biomarker of prostate cancer malignancy [105,112]. Therefore, Western blotting was performed on the proteins extracted from ~1x10⁷ cells from each line to identify total Hsp-27 as well as the differential phosphorylation of this protein at Ser15, Ser78 and Ser82. All methodologies used were identical to those previously reported [33].

Glycoconjugate expression. The behavior of malignant epithelial cells is influenced by expression of cell-surface complex glycoconjugates, particularly sialic acids [46,113–115]. To assess the potential effects of RPL19 knockdown on cell-surface oligosaccharide expression, cell-blocks were prepared from cell-lines PNT2, PC-3Mparental, PC-3Mscramble, and si-RPL19-PC-3Mtrue ST-3. Cell pellets were processed and embedded in paraffin wax blocks [33,116]. Sections were cut at 5 μm and stained for Neu5Acα2–→3Gal- and Neu5Acα2–→6Gal- using biotinylated lectins (Vector Laboratories, Peterborough, UK) from Sambucus nigra and Maackia amurensis respectively [117,118]. The biotinylated lectins from Ulex europaeus, Lotus tetragonolobus and Aleuria aurantia were employed to detect terminal fucosyl linkages. Negative controls included the absence of staining when the lectins were not included in the staining protocol and the abolition of staining following pre-treatment of the slides with neuraminidase prior to incubation with the lectins [119]. Lectin-binding was detected using an avidin-peroxidase conjugate visualized following polymerization of 3,3’-diaminobenzidine (DAB).

Supporting Information Material

This information contains Supporting Information Figure S1 confirming the validation of the arrays by PCR and additional data-tables providing detailed information on the alterations in gene expression, including their involved networks, induced following knockdown of RPL19. This material supports, but does not extend, the findings and conclusions of this study.

Supporting Information

Figure S1 As well as the gene-sequences employed previously [33] four additional sequences were used to interrogate genes up-regulated and down-regulated and thus validate the levels of expression detected by array-analysis. As with the arrays, the levels were quantified relative to PC-3Mscramble cells that were set at unity. (TIFF)

Table S1 Top 50 genes up-regulated (fold change). Genes are arranged in descending order according to log₂ fold change with corresponding p-values.

| Gene | Fold Change | p-value |
|------|-------------|---------|
|     |             |         |

Table S2 Top 50 genes down-regulated (fold change). Genes are arranged in descending order according to log₂ fold change with corresponding p-values.

| Gene | Fold Change | p-value |
|------|-------------|---------|
|     |             |         |

Table S3 Gene ontology terms. Gene ontology (GO) biological process terms found to be significantly associated with genes significantly differentially expressed after knockdown of RPL19 using hypergeometric tests.

Table S4 KEGG pathways. KEGG pathways containing genes significantly differentially expressed after RPL19 knockdown using hypergeometric tests.

Table S5 Pathways modulated after RPL19 knockdown. Top five interlinked pathways containing genes significantly differentially expressed after RPL19 knockdown using hypergeometric tests.

Table S6 Gene ontology molecular function terms. Gene ontology (GO) molecular function terms significantly associated with genes differentially expressed after knockdown of RPL19 using hypergeometric tests.

Table S7 Glycosyltransferase and ion-channel genes modulated following RPL19 knockdown. Following RPL19 knockdown, modulated expression of only two glycosyltransferase genes was detected but with more profound changes to ion channels indicating significant changes to cellular homeostasis.

Table S8 Characteristics of antibodies used to analyze changes in proteins expressed following RPL19 knockdown. Details of protein expression by Western Blotting analysed using a range of mono-specific antibodies to define changes in cellular phenotype following RPL19 knockdown.

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Author Contributions

Conceived and designed the experiments: AB CSF. Performed the experiments: GB AD SF TD PG SY. Analyzed the data: DB BL MBAD CMG CSC. Contributed reagents/materials/analysis tools: CSC MBAD YK. Wrote the paper: AB CSF CSC MBAD.

References

1. Gough J, Karplus K, Hughey R, Chothia C (2001) Assignment of homology to genome sequences using a library of hidden Markov models that represent all proteins of known structure. Journal of Molecular Biology 313: 903–919.
2. Delaunay J, Schapira G (1974) Phylogenic distance between prokaryotes and eukaryotes as evaluated by ribosomal proteins. FEBS Letters 40: 97–100.
3. Wittmann HG (1980) Structure and evolution of ribosomes. Molecular Biology, Biochemistry and Biophysics 32: 376–397.
4. Taylor DJ, Devkota B, Huang AD, Topf M, Narayanan E, et al. (2009) Comprehensive molecular structural structure of the eukaryotic ribosome. Structure 17: 1591–1604.
5. Warner JR, McIntosh KB (2009) How common are extraribosomal functions of ribosomal proteins. Molecular Cell 34: 3–11.
6. Weinberg RA (2008) Transcription by moonlight: structural basis of an extraribosomal activity of ribosomal protein S10. Molecules and Cells 32: 747–748.
7. Marechal V, Elenbaas B, Piette J, Nicolas JC, Levine AJ (1994) The ribosomal L5 protein is associated with mdm-2 and mdm-2-p53 complexes. Molecular and Cellular Biology 14: 7414–7420.
8. Dai MS, Arnold H, Sun XX, Sears R, Lu H (2007) Inhibition of c-Myc activity by ribosomal protein L11. EMBO Journal 26: 3332–3345.
9. Chauhuri S, Vyas K, Kapasi P, Komar AA, Dinman JD, et al. (2007) Human ribosomal protein L13A is dispensable for canonical ribosome function but indispensable for efficient rRNA methylation. RNA 13: 2224–2237.
10. Chen D, Zhang B, Li M, Wang W, Li Y, et al. (2007) Ribosomal Protein S7 as a novel immodulator of p53-MDM2 interaction: binding to MDM2 stabilization of p53 protein and activation of p53 function. Oncogene 26: 5029–5037.
11. Lai K, Amsterdam A, Farrington S, Bronson RT, Hopkins N, et al. (2009) Many ribosomal protein mutations are associated with growth impairment and tumor predisposition in zebrafish. Developmental Dynamics 238: 76–85.
12. MacInnes AW, Amsterdam A, Whitaker CA, Hopkins N, Gees JA (2008) Loss of p53 synthesis in zebrafish tumors with ribosomal protein gene mutations. Proceedings of the National Academy of Science USA 105: 10408–10413.

13. Luft F (2010) The rise of a ribosomalopathy and increased cancer risk. Journal of Molecular Medicine 88: 1–3.

14. Ellis SR, Gieles PE (2011) Diamond Blackfan anemia: ribosomal proteins going rogue. Seminars in Hematology 48: 89–96.

15. Campagnoli MF, Ramegh U, Arminghio M, Quarello P, Garelli E, et al. (2008) RPS19 mutations in patients with Diamond-Blackfan anemia. Human Mutation. 29: 911–920.

16. Dutt S, Narla A, Lin K, Mullally A, Abayasekara N, et al. (2011) Riboflavin from patients with Diamond-Blackfan anemia show abnormal expression of genes involved in protein synthesis, amino acid metabolism and cancer. BMC Genomics 10: 442.

17. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

18. Dutt S, Narla A, Lin K, Mullally A, Abayasekara N, et al. (2011) Riboflavin from patients with Diamond-Blackfan anemia show abnormal expression of genes involved in protein synthesis, amino acid metabolism and cancer. BMC Genomics 10: 442.

19. Ellis SR, Gieles PE (2011) Diamond Blackfan anemia: ribosomal proteins going rogue. Seminars in Hematology 48: 89–96.

20. Yang D, Kusser I, Kopke AK, Koop BF, Matheson AT (1999) The structure and evolution of mammalian ribosomal proteins. Biochemistry and Cell Biology 73: 933–947.

21. Yang D, Kasper I, Kopke AK, Koop BF, Matheson AT (1999) The structure and evolution of the ribosomal proteins encoded in the spc operon of the archaeon (Crenarchaeota). Salibfusis acidocaldarius. Molecular Phylogenetics and Evolution 12: 177–185.

22. Chen YL, Lin A, McNally J, Peleg D, Meyahu O, et al. (1987) The primary structure of rat ribosomal protein L19. A determination from the sequence of nucleotides in cDNA and from the sequence of amino acids in the protein. Journal of Biological Chemistry 262: 1111–1115.

23. Dressman MA, Baras A, Malinowski R, Alvis LB, Kwon I, et al. (2003) Gene expression profiling detects gene amplification and differentiates tumor types in breast cancer. Cancer Research 63: 2194–2199.

24. Kauraniemi P, Himmelstien A (2006) Activation of multiple cancer-associated genes by the ERBB2 amplification in breast cancer. Endocrine-Related Cancer 13: 39–49.

25. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

26. Minner S, Jesen B, Stiererth L, Burundi E, Kollermann, J, et al. (2010) Low-level HER2 overexpression is associated with rapid tumor cell proliferation and poor prognosis in prostate cancer. Clinical Cancer Research 16: 1553–1560.

27. Ransome MT, Muriro K, Betti C, Pica E, Zottl G, et al. (2010) Detection of HER2 amplification using the SISH technique in breast, colon, prostate, lung and ovarian carcinoma. Anticancer Research 30: 1287–1292.

28. Huang CJ, Chien CC, Yang SHC, Chien CC, Sun HL, Cheng YC, et al. (2008) Faecal ribosomal protein L19 is a genetic prognostic factor for survival in colorectal cancer. Journal of Cellular and Molecular Medicine 12: 1936–1943.

29. Svaren J, Ehrig T, Abdulkadir SA, Ehrengruber MUW, M A, Milbrandt J (2003) A novel function of differentiation revealed by cDNA microarray analysis. Journal of Biological Chemistry 278: 3072–3078.

30. Uyama T, Kitagawa H, Tanaka J, Tamura J, Ogawa T, et al. (2003) Eosinophyl and ribosomal proteins encoded in the spc operon of the archaeon (Crenarchaeota). Salibfusis acidocaldarius. Molecular Phylogenetics and Evolution 12: 177–185.

31. Uyama T, Kusser I, Kopke AK, Koop BF, Matheson AT (1999) The structure and evolution of ribosomal proteins encoded in the spc operon of the archaeon (Crenarchaeota). Salibfusis acidocaldarius. Molecular Phylogenetics and Evolution 12: 177–185.

32. Chan YL, Lin A, McNally J, Peleg D, Meyahu O, et al. (1987) The primary structure of rat ribosomal protein L19. A determination from the sequence of nucleotides in cDNA and from the sequence of amino acids in the protein. Journal of Biological Chemistry 262: 1111–1115.

33. Dressman MA, Baras A, Malinowski R, Alvis LB, Kwon I, et al. (2003) Gene expression profiling detects gene amplification and differentiates tumor types in breast cancer. Cancer Research 63: 2194–2199.

34. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

35. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

36. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

37. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

38. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

39. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

40. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

41. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

42. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

43. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

44. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

45. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

46. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

47. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

48. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.

49. Mano MS, Rosa DD, De Azambuja E, Ismael GFV, Durbecq V (2007) The structure, assembly and evolution of the complete set of 60s ribosomal proteins and their expression. Marine Biotechnology 9: 336–381.
66. Ruggero D, Pandolfi PP (2003) Does the ribosome translate cancer? Nature Reviews Cancer 3: 179–192.

67. Yang SH, Huang CJ, Lee CL, Lai CC, Chien CC, et al. (2010) Fetal RNA detection of cytokeratin 19 and ribosomal protein L19 for colorectal cancer.

68. Vaarala MH, Porvari KS, Kyllonen AP, Mustonen MV, Lukkarinen O, et al. (1998) Several genes encoding ribosomal proteins are over-expressed in prostate-cancer cell lines: confirmation of L7a and L37 over-expressed in prostate cancer tissue samples. International Journal of Cancer 78: 27–32.

69. Henry JL, Coggin DL, King CR (1993) High-level expression of the ribosomal protein L19 in human breast tumors that overexpress erb-B2. Cancer Research 53: 1409–1408.

70. Hatakeyama T, Kaufmann F, Schroeter B, Hatakeyama T (1989) Primary structures of five ribosomal proteins from the archaeabacterium Halo bacterium marismortui and their structural relationships to eubacterial and eukaryotic ribosomal proteins. European Journal of Biochemistry 183: 683–693.

71. Bardet M, Cramoisyvski S, Grabisuk B, Bogert I, Panic I, et al. (2009) The p53 tumor suppressor causes congenital malformations in Rpl24-deficient embryos and promotes their survival. Molecular and Cellular Biology 29: 2419–2404.

72. Zhao C, Zhang D, Jin Y, Wu H, Liu Z, et al. (2011) Mutation in ribosomal protein L21 underlies hereditary hypotrichosis simplex. Human Mutation March. pp 15 (Epub ahead of print).

73. Brachveg B, Dikчасas J, Mohr H, Weizel H, von der Mark K, et al. (2003) Annexin A3 is not essential for skeletal development. Molecular and Cellular Biology 23: 2097–2111.

74. Jonkers J, van Amerongen R, van der Valk M, Reuben-Maandag E, Molenaar M, et al. (1999) In vivo analysis of Fra1 deficiency suggests a compensatory role of Fra3. Mechanisms of Development 88: 183–191.

75. Leong WF, Chau JF, Li B (2009) p53 deficiency leads to compensatory up-regulation of p16INK4a. Molecular Cancer Research 7: 354–360.

76. Belin S, Beghin A, Solano-Gonzalez E, Bezin L, Brunet-Manquat S, et al. (2009) Regulation of ribosome biogenesis and translational capacity is associated with tumor progression of human breast cancer cells. PLoS One 4: e7147.

77. Ke S, Zhang XH, Chasin LA (2008) Positive selection acting on splicing motifs reflects compensatory evolution. Genome Research 18: 533–543.

78. Custodio N, Carmona-Fonseca M (2001) Quality control of gene expression in the nucleus. Journal of Cellular and Molecular Medicine 5: 267–275.

79. Mauzner B, Bampath P, Seshadri V, Maitra RR, DiCorleto PE, et al. (2003) Heat shock protein (HSP) expression independently predicts clinical outcome in prostate cancer. American Journal of Pathology 154: 137–144.

80. Foster CS, Dodson AR, Ambrose L, Fisher G, Moller H, et al. (2009) HSP-70 expression at diagnosis predicts poor clinical outcome in prostate cancer independent of ETS-gene rearrangement. British Journal of Cancer 101: 1137–1144.

81. Beerenwinkel N, Pachter L, Sturmfels B, Elena SF, Lenski RE (2007) Analysis of genome-wide single nucleotide polymorphisms on protein-protein interactions. Biophysical Journal 96: 2178–2180.

82. Benjamini Y, Hockberg Y (1995) Controlling the false discovery rate: A practical and powerful approach to multiple testing. Journal of the Statistical Society Series B 57: 289–300.

83. Bell G, Gentleman RC (2007) Bioconductor: open software development for computational biology and bioinformatics. Genome Biology 5: R80.

84. Smyth G (2004) Linear models and empirical bayes methods for assessing differential expression in microarray experiments. Statistical Applications in Genetics and Molecular Biology 3: Article 3. Epub 2004 Feb 12.

85. Ritchie ME, Silver J, Oshlack A, Holmes M, Divagama D, et al. (2007) A comparison of background correction methods for two-colour microarrays. Bioinformatics 23: 2700–2707.

86. Cornford P, Evans J, Dodson AR, Parsons K, Woolfenden M, et al. (1999) Protein kinase C isoenzyme patterns characteristically modulated in early prostate cancer. American Journal of Pathology 154: 137–144.

87. Strom GD, Speed T (2003) Normalization of cDNA microarray data. Methods 31: 265–273.

88. Huffman LJ, Stojadinovic A, Nicholson GL (1984) Cell surface properties of spontaneously immortalized cells of the rat mammary adenocarcinoma. Cancer Research 44: 527–532.

89. Dolger D, Saurheen H, Slabbert H, Hopwood J, Woolfenden K, et al. (2003) Putting a finger on growth pathways in the metastatic process. BMC Cancer 7: 64.

90. LaTulippe E, Satapojjan J, Smith A, Scher H, Scardino P, et al. (2002) Comprehensive gene expression analysis of prostate cancer reveals distinct transcriptional programs associated with metastatic disease. Cancer Research 62: 4499–4506.

91. Casamayor O, Benito P, Berger R, Mosowiczcz I, Faull E, et al. (1991) Immortalization of human adult normal prostatic epithelial cells by liposomes containing large T-SV40 gene. Journal of Urology 146: 881–886.

92. Kajihara ME, Shankar N, Ohnuki Y, Lechner JF, Jones LW (1979) Establishment and characterization of a human prostate carcinoma cell line (PC-3). Investigative Urology 17: 16–23.

93. Shevlin DH, Gorny KI, Kukreja SC (1989) Patterns of metastasis by the human prostate cancer cell line PC-3 in athymic nude mice. Prostate 15: 167–194.

94. Smith P, Rhodes NP, Belsey C, Ke Y, Foster CS (1998) Prostatic stromal cell phenotype is directly modulated by neuropilin-1. Urology 51: 663–670.

95. Solier S, Barb J, Zeeberg BR, Varma S, Ryan M, et al. (2010) Genome-wide analysis of novel splice variants induced by topoisomerase I poisoning shows preferential occurrence in genes encoding splicing factors. Cancer Research 70: 8055–8065.

96. Cussenot O, Berthon P, Berger R, Mosowiczcz I, Faull E, et al. (1991) A scoring system for immunohistochemical staining: consensus report of the task force for the research of the EORTC-GCCG. Journal of Clinical Pathology 50: 801–804.

97. Cornwall P, Evans J, Dodson AR, Parsons K, Wolffenbrouck K, et al. (1999) Protein kinase C isoenzyme patterns characteristically modulated in early prostate cancer. American Journal of Pathology 154: 137–144.

98. Foster CS, Dodson AR, Ambrose L, Fisher G, Moller H, et al. (2009) HSP-70 expression at diagnosis predicts poor clinical outcome in prostate cancer independent of ETS-gene rearrangement. British Journal of Cancer 101: 1137–1144.

99. Benjamini Y, Yekhlansky S, Holmes M, Divagama D, et al. (2007) A comparison of background correction methods for two-colour microarrays. Bioinformatics 23: 2700–2707.

100. Smyth G (2004) Linear models and empirical bayes methods for assessing differential expression in microarray experiments. Statistical Applications in Genetics and Molecular Biology 3: Article 3. Epub 2004 Feb 12.

101. Benjamini Y, Yekhlansky S, Holmes M, Divagama D, et al. (2007) A comparison of background correction methods for two-colour microarrays. Bioinformatics 23: 2700–2707.

102. Steck PA, Nicolson GL (1984) Cell surface properties of spontaneously immortalized cells of the rat mammary adenocarcinoma. Cancer Research 44: 527–532.

103. de Albuquerque Garcia Redondo P, Nakamura CV, de Souza W, Morgado-Jesus S, Camargo E, et al. (2003) Changes in cell-surface oligosaccharides after growth of the human prostate cancer cell line PC-3 in athymic nude mice. Hepatogastroenterology 50: 710–715.

104. Cornford P, Evans J, Dodson AR, Parsons K, Woolfenden M, et al. (1999) Protein kinase C isoenzyme patterns characteristically modulated in early prostate cancer. American Journal of Pathology 154: 137–144.

105. Ritchie ME, Silver J, Oshlack A, Holmes M, Divagama D, et al. (2007) A comparison of background correction methods for two-colour microarrays. Bioinformatics 23: 2700–2707.

106. Ritchie ME, Silver J, Oshlack A, Holmes M, Divagama D, et al. (2007) A comparison of background correction methods for two-colour microarrays. Bioinformatics 23: 2700–2707.