Telomere length in the two extremes of abnormal fetal growth and the programming effect of maternal arterial hypertension

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We tested the hypothesis that leukocyte telomere length (LTL) is associated with birth weight in both extremes of abnormal fetal growth: small (SGA) and large for gestational age newborns (LGA). Clinical and laboratory variables of the mothers and the neonates were explored; 45 newborns with appropriate weight for gestational age (AGA), 12 SGA and 12 LGA were included. Whether the differences might be explained by variation in \textit{OBFC1} (rs9419958) and \textit{CTC1} (rs3027234) genes associated with LTL was determined. A significant association between birth weight and LTL was observed; LTL was significantly shorter in LGA newborns (1.01 ± 0.12) compared with SGA (1.73 ± 0.19) \(p < 0.005\), mean ± SE. Maternal (Spearman \(R = 0.6, p = 0.03\)) and neonatal LTL (\(R = 0.25, p = 0.03\)) were significantly and inversely correlated with maternal history of arterial hypertension in previous gestations. Neonatal LTL was not significantly associated with either rs9419950 or rs3027234, suggesting that the association between neonatal LTL and birth weight is not influenced by genetic variation in genes that modify the interindividual LTL. In conclusion, telomere biology seems to be modulated by abnormal fetal growth; modifications in telomere length might be programmed by an adverse environment in utero.

The thrifty phenotype/genotype hypothesis for explaining the origin of metabolic syndrome-associated phenotypes was introduced several years ago. Since the first description of David Barker and coworkers about the fetal metabolic programming of adult chronic diseases, there has been a large amount of epidemiological data confirming the strong association between an impaired intrauterine environment and adult metabolic and cardiovascular diseases (CVD).

The concept of fetal programming has two important emerging putative consequences: the transmission of the phenotype from the mother to the progeny and even through generations, and the permanent change of the metabolism of the newborn exposed to an adverse intrauterine environment even in the absence of the original stimulus. In line with the last point, all the robust epidemiological data supporting the concept of fetal programming was specifically focused on birth weight and its relation with adult chronic diseases. In this regard, the most suitable hypothesis to explain the interplay between the in utero environment and the later development of chronic diseases is the concept of epigenetic modifications, which illustrates how the in utero life modulates the adult phenotype and suggests the induction of permanent genomic marks. For instance, some interesting evidence from experimental studies showed that twinning and undernutrition in the mother around the time of conception alter the epigenetic status of genes involved in the regulation of metabolism in hypothalamus. Specifically, an adverse in utero environment is associated with epigenetic changes in the methylation status of proopiomelanocortin and the glucocorticoid receptor (GR) that might have an impact on the regulation of food intake in offspring when they reach adulthood, leading to the development of obesity.
In line with these data, Lillycrop and coworkers showed that feeding a protein-restricted diet to pregnant rats causes hypomethylation of GR in umbilical cord and a reduced Dnmt1 expression\(^2\). Moreover, we demonstrated that maternal diet during pregnancy has an impact on fetal metabolic programming through changes in liver mitochondrial DNA content and liver transcriptional activity of the peroxisome proliferator-activated receptor gamma, coactivator (Ppargc1a)\(^\dagger\).

Furthermore, evidence from human studies examining changes in the global methylation status of cord blood DNA identified differential methylation at 19 cytosine-guanine dinucleotides associated with either decreased or increased birth weight\(^\dagger\). We also reported that changes in methylation of genes related with mitochondrial biogenesis such as PPARGCIA is associated with small and large in comparison with normal for gestational age\(^\dagger\). Finally, remarkable findings about offspring exposed to gestational diabetes mellitus support the concept of fetal programming of adult metabolic diseases\(^\dagger\).

Recent evidence suggests that telomere biology is potentially plausible of modification during fetal life\(^\dagger\), and modifications in Dnmt1 of GR in umbilical cord and a reduced ing a protein-restricted diet to pregnant rats causes hypomethylation potential methylation at 19 cytosine-guanine dinucleotides associated the global methylation status of cord blood DNA identified differ-

### Results

Clinical features, anthropometrical variables, and laboratory findings of the pregnant women (at the time of delivery) and the neonates according to the offspring birth weight can be seen in Table 1. Many features of the mother and newborn associated with birth weight are consistent with those already described\(^\dagger\), that is, pregestational body mass index (BMI) of the mother is associated with neonatal sex and gestational age-adjusted body weight. As expected, newborn leptin correlated with neonatal sex and gestational age-adjusted body weight.

Interestingly, we observed a significant association between sex and gestational age-adjusted birth weight and LTL (Figure 1A); LTL was significantly shorter in the LGA newborns than in the SGA ones. Accordingly, LTL was significantly and inversely correlated (Spearman R: \(-0.21, p=0.008\)) with newborn’s body weight Z-score (Figure 1B). In addition, LTL was significantly and inversely associated with other developmental characteristics of the newborns, such as neonatal head circumference (Spearman correlation R = \(-0.3, p = 0.03\)). There were no sex differences in LTL among the neonates.

We also explored the LTL in the mothers and observed that there was no statistically significant association with the newborns birth weight (Figure 2A). Nevertheless, we found that maternal LTL was significantly and inversely correlated with history of arterial hypertension in previous pregnancies (Spearman R = \(-0.6, p = 0.03\)); differences in mothers’ LTL according to history of elevated arterial blood pressure in previous pregnancies are shown in Figure 2B. Significant differences in mothers’ systolic and diastolic blood pressure according to previous history of arterial hypertension were observed and are shown in Supplementary Figure 1.

Neonatal LTL was also significantly and inversely correlated with maternal history of arterial hypertension in previous pregnancies (Spearman R = \(-0.25, p = 0.03\)) and significantly influenced by the arterial blood pressure status of their mothers in current pregnancy (Figure 2C). These findings are not surprising because of the association of hypertension in successive pregnancies: 33% of mothers with previous antecedents of hypertension developed

### Table 1 | Clinical features, anthropometric variables, and laboratory findings of the pregnant women and the newborns according to the neonates’ birth weight

|                        | SGA          | AGA          | LGA          | P level: SGA vs AGA, SGA vs LGA, AGA vs LGA* |
|------------------------|--------------|--------------|--------------|---------------------------------------------|
| Number of new born     | 6            | 45           | 12           |                                             |
| Birth weight adjusted  | \(-1.840.21\)| 0.20 ± 0.12  | 1.67 ± 0.09  | 0.00001, 0.00008, 0.000001                   |
| and sex. Z score       |              |              |              |                                             |
| Neonate birth weight, g| 2435 ± 114.5 | 3387.4 ± 126.9 | 4311 ± 52 | 0.0001, 0.0003, 0.00001                     |
| Head circumference, cm | 32.3 ± 0.43  | 34.3 ± 0.4   | 36.1 ± 0.39  | 0.002, 0.0002, 0.004                        |
| Neonate length, cm     | 46 ± 0.74    | 48.1 ± 0.78  | 51.8 ± 0.61  | 0.007, 0.003, 0.003                        |
| Apgar score            | 8.3 ± 0.36   | 8.14 ± 0.34  | 9 ± 0        | NS, NS, NS                                  |
| Gestational age, weeks | 38.0 ± 0.48  | 38.2 ± 0.42  | 39.8 ± 0.21  | NS, 0.006, 0.02                            |
| Leptin, ng/mL          | 7.07 ± 1.42  | 16.1 ± 2.05  | 27.0 ± 4.56  | 0.025, 0.0003, 0.02                        |
| Plasma homocysteine    | 10.82 ± 1.6  | 9.26 ± 0.65  | 8.75 ± 0.70  | NS, NS, NS                                  |
| levels                 |              |              |              |                                             |

### Clinical features and anthropometric variables of pregnant women at the time of delivery according to the offspring’s birth weight

|                        | SGA          | AGA          | LGA          |                                     |
|------------------------|--------------|--------------|--------------|-----------------------------------|
| Age, years             | 28.3 ± 1.8   | 31.8 ± 1.9   | 28.0 ± 2.3   | NS, NS, NS                        |
| Physical activity, hours/week | 0.66 ± 0.6 | 1.0 ± 0.37   | 0.0 ± 0.0    | NS, NS, NS                        |
| Number cigarettes/day  | 2.5 ± 1.3    | 2.9 ± 1.3    | 0.91 ± 0.8   | NS, NS, NS                        |
| Pregestational BMI, kg/m² | 21.9 ± 2.2  | 22.9 ± 0.7   | 26.6 ± 1.5   | NS, 0.03, 0.03                    |
| Body weight gain during pregnancy, Kg | 12.8 ± 1.4 | 12.0 ± 1.0   | 15.3 ± 1.0   | NS, NS, NS                        |
| DABP, mmHg             | 70.0 ± 2.65  | 71.7 ± 1.8   | 72.1 ± 2.2   | NS, NS, NS                        |
| SABP, mmHg             | 115.2 ± 3.0  | 114.1 ± 1.9  | 115.8 ± 2.68 | NS, NS, NS                        |

SGA: small for gestational age. AGA: appropriate for gestational age. LGA: large for gestational age. Results are expressed as mean ± SE. P stands for the level of statistical significance by Mann-Whitney test. NS: not significant.
hypertension in the current pregnancy against 5% in those who do not (chi square, $p < 0.02$).

**Neonatal LTL and gene variants.** Because previous genome-wide association studies (GWASs) have identified some single nucleotide polymorphisms (SNPs) associated with LTL variation, such as the rs9419958 at *OBFC1* oligonucleotide/oligosaccharide-binding fold locus and the rs3027234 at *CTC1* telomere maintenance complex locus, we decided to explore whether these genetic variants accounted for the observed differences in the neonatal LTL.

Figure 1 | (A)—Neonatal leukocyte telomere length (LTL)/ single-copy gene (HBB, human β-globin) ratio in small for gestational age (SGA), appropriate for gestational age (AGA) and large for gestational age (LGA) neonates. P values stands for the post hoc Newman-Keuls test after ANOVA with log-transformed variables. (B)—Pearson correlation test of log transformed values of LTL sex and and age-adjusted neonatal birth weight. R: $-0.344, p=0.0046$.

Previous reports support the selection of these loci for exploration in our sample. For example, variation in *OBFC1* showed a replicated association with mean LTL in a large scale GWA meta-analysis of 37,684 individuals from 15 cohorts. In addition, a large population study on the risk of CVD showed that the rs9419958 was associated with CVD-related death, particularly among women. The *OBFC1* gene also appears to function in a telomere-associated complex of proteins required to protect telomeres from DNA degradation. Finally, functional experiments demonstrated that *OBFC1* represents a new player in the telomere interactome for telomere maintenance.

Figure 2 | (A)—Mother LTL/single-copy gene (HBB, human β-globin) ratio according to birth weight (small for gestational age (SGA), appropriate for gestational age (AGA), and large for gestational age (LGA). (B)—Mother LTL/single-copy gene (HBB, human β-globin) ratio according to their previous history of arterial hypertension. (C)—Neonatal LTL/single-copy gene (HBB, human β-globin) ratio according to the current status of maternal blood pressure.
Regarding to CTC1 gene, there is biological evidence that supports the selection of this locus for exploration of genetic variants. For instance, this gene encodes a component of the CST complex, which plays an essential role in protecting telomeres from degradation. Furthermore, CTC1 may have a function in DNA metabolism and in regulating telomere homeostasis; the CST complex seems also to play critical roles in coordinating telomerase elongation and fill-in synthesis to complete telomere replication.

Allelic frequencies and genotype distributions of the rs9419958 and rs3027234 were in Hardy-Weinberg equilibrium. For the analysis, we used a recessive model of inheritance because of the low frequency of the minor allele of each SNP in our sample.

We observed that the neonatal LTL was not significantly associated with either rs9419950 or rs3027234 (Table 2), suggesting that the association between neonatal LTL and birth weight is not influenced by genetic variation in genes that modify the interindividual LTL (Table 2). The rs9419958 was significantly correlated with neonatal plasma homocysteine levels (Spearman R = 0.26, p < 0.02 and Table 2) and significantly and negatively correlated with the mother’s systolic arterial blood pressure (Spearman R = -0.24, p < 0.04).

To better understand these unexpected findings associated with the gene variants, we performed a functional association analysis of interactions pathways of OBFC1 and CTC1 genes and its relation with arterial hypertension and homocysteine levels allowing the exploration of disease-gene interactions. The functional analysis showed a predicted disease-gene node around OBFC1 but not CTC1 that integrates an interrelated meta-node composed of two genes/proteins: 1) HNF1 homeobox a (alias TCF1)—a transcription factor that is strongly associated with the angiotensinogen (AGT), which is associated with hypertension, and 2) HNF4A (hepatocyte nuclear factor 4 alpha) that is connected with the methylenetetrahydrofolate dehydrogenase (MTHFD1) and was predicted to modulate homocysteine levels (Figure 3).

Discussion
In this study, we asked whether newborn telomere biology might be modulated by abnormal fetal growth and observed that birth weight was significantly associated with LTL. Our results show for the first time that babies who are born large have shorter telomeres in comparison with small babies. Accordingly, LTL was also significantly and inversely correlated with anthropometric variables associated with newborn’s body size, such as head circumference. Consistent with previous human reports, LTL of SGA babies was not significantly different from that of AGA babies, suggesting that the molecular mechanisms behind decreased fetal growth are unrelated to telomere biology. A note of caution clarifying that the term of SGA is not synonymous with “intrauterine growth retardation (restriction)” must be added.

We might wonder about the factors that modulate the telomere biology in larger babies with shorter telomeres. LGA and SGA babies face many challenges during fetal life, but the reason the LGA babies are different in terms of telomere biology remains largely unknown. Nevertheless, telomere biology might be suggested as an underlying mechanism connecting fetal programming and chronic adult diseases, as the telomere homeostasis is quite receptive of the in utero environment and is also highly sensitive to inflammation, oxidative stress, and related adverse and stressful situations. Although homocysteine levels may be one of them, we were not able to find any significant association.

Moreover, LGA infants are well known to suffer from some complications, including fetal hypoxia and hypoglycemia, which is a consequence of mother hyperinsulinemia. Still, whether these or other factors might be the determining ones of the modifications of the telomere length in LGA babies remains to be explored. In agreement with the hypothesis of the in utero reprogramming toward senescence, previous evidence showed higher cord blood telomerase activity in Type 1 and gestational diabetes.

Furthermore, when we explored whether maternal factors might be able to modify babies’ LTL at birth, we observed that arterial hypertension was a critical factor in the modulation of the newborn LTL. We found that LTL at birth was significantly associated not only with arterial blood pressure of their mothers during current pregnancy but also with the history of arterial hypertension in previous pregnancies, despite the fact that most mothers who developed hypertension in the current pregnancy were devoid of personal history of hypertension. This finding supports the notion about the maternal disease-associated programming effect of the newborn telomere length. Although this finding is expected because CVDs, such as hypertension, are associated with short LTL, the programming effect of the mother’s hypertension is a novel finding and suggests that the arterial blood pressure control is an important goal to achieve in women during fertile age, although prospective studies are necessary to further investigate this issue.

On the other hand, we asked whether genetic factors might influence LTL at birth and then explored two SNPs in genes associated either with telomere biology (OBFC1) or with LTL (OBFC1 and CTC1). Interestingly, we observed that genetic variation does not account for LTL at birth. However, an unexpected result that needs further confirmation, such as the significant association between the rs9419958 and fetal plasma homocysteine levels and maternal systolic blood pressure, was observed. Functional association analysis supports the biological plausibility of our findings; specifically, it was shown that OBFC1 interacts with HNF4A as explained in the results.
section. Notably, a GWAS in the community-based Framingham Heart Study on risk factors of CVD showed an association between a variant in \textit{OBFC1} gene (the rs3814219) and flow-mediated dilation, a surrogate of impaired endothelial function\(^{36}\); this variant was among the top SNPs associated with brachial artery vascular function-related traits (\(p\) value for association 9.48 \(\times 10^{-7}\))\(^{36}\). These results were replicated in a large sample of subjects from Europe, showing that variants located in \textit{OBFC1} are associated with higher risk of CVD, including coronary heart disease\(^{37}\); remarkably, higher expression of \textit{OBFC1} in adventitial aortic tissue was demonstrated in this study\(^{37}\). Taken together, these findings support our results and suggest a strong biological plausibility of the association between the \textit{OBFC1} locus and the risk of CVD in humans.

The association between the variant in \textit{OBFC1} locus and fetal plasma homocysteine levels might represent a CVD-risk related-associated trait in consonance with the association found with maternal systolic blood pressure, as it was previously shown that there is a strong linear association between the maternal and fetal levels of homocysteine in circulation\(^{38}\) and that elevated homocysteine levels are a strong and independent risk factor for vascular disease and hypertension in adult\(^{39}\) and adolescents\(^{40}\).

Finally, we would like to discuss the putative clinical implications of our results. Although the design of our study prevents against any further speculation about adult health outcomes of the studied babies, some concluding remarks might be added. For instance, although long-term consequences of abnormal fetal growth are already known, specifically those related with LGA infants who are at increased risk of obesity, type 2 diabetes, and CVD in adolescence and adulthood\(^{4}\), some novel data emerged. Our findings might suggest a "programming senescence effect" of adult chronic diseases associated with the metabolic syndrome, particularly arterial hypertension. As neonatal peripheral blood LTL is shown to be highly correlated with telomere length in tissues, we might speculate that both abnormal fetal growth and maternal stress have an impact on offspring telomere biology, leading to a "short-telomere phenotype" that renders them prone to cardiovascular complications in adulthood. This shorter telomere phenotype at birth indeed is imprinting an accelerated aging phenotype in target tissues that might

![Figure 3](image)

**Figure 3** Analysis of interaction pathways of \textit{OBFC1} and \textit{CTC1} genes (pink nodes) and its relation with arterial hypertension and homocysteine levels. The integrative analysis of biological networks was performed using "VisANT 4.0: Integrative network platform to connect genes, drugs, diseases, and therapies" (a freely available and Web-based workbench at http://visant.bu.edu)\(^{47}\).
compromise cellular function and integrity, affecting normal physiology and increasing disease risk.

A limitation of our study should be highlighted as a larger-sized sample and availability longitudinal clinical and biochemical measurements in the infants would considerable strengthen our investigation. In addition, we could have limited statistical power to detect modest genetic effects, given the small sample size of our sample; finally, the extrapolation of our genetic findings to other ethnicities remains to be explored.

In conclusion, in this study, we tested the hypothesis that leukocyte telomere length (TLT) may be associated with birth weight in both extremes of abnormal fetal growth: small for gestational age and large for gestational age newborns. Our results show that babies who are born large have shorter telomeres in comparison with small babies. Furthermore, when we explored whether maternal factors might be able to modify babies’ TLT at birth, we observed that arterial hypertension was a critical factor in the modulation of the newborn TLT. Our findings might suggest a “programming senescence effect” of adult chronic diseases associated with the metabolic syndrome, particularly arterial hypertension.

Methods

Ethics statements. This study was performed according to the principles of the Declaration of Helsinki and was approved by the institutional review board and the bioethical committee of our institution. Furthermore, written consent was obtained from all the patients.

Subjects. Sixty-nine consecutive healthy pregnant women and their babies were included in this study; selecting around two AGA babies for each SGA or LGA groups. According to the offspring birth weight normalized by sex and gestational age, there were 45 newborns with appropriate weight for gestational age (AGA) and 24 with abnormal weight for gestational age: 12 SGA and 12 LGA.

There were no multiple pregnancies in the study population. None of the mothers had either preeclampsia-eclampsia or gestational diabetes or were under any medication. At the time of delivery, an aliquot of cord blood was stored for further analysis.

All the investigations performed in this study were conducted in accordance with the guidelines of the Declaration of Helsinki. Written consent from the patients had been granted, in accordance with the procedures approved by the ethical committee of our institution. Complete medical, obstetrical, and perinatal data were recorded as shown below.

Obstetric Variables. Gestational age was defined on the basis of the last menstrual period, physical examination, and ultrasonographic biometry. Length of gestation at delivery was measured as the number of weeks of gestation. The ultrasound measure during the first trimester took precedence if these measures did not agree in terms of dating gestational age at birth.

Maternity data including age, geographical origin, pregestational BMI, education, smoking status (number of cigarettes/day), history of abortion, history of preterm birth and previous abnormal fetal weight, preeclampsia-eclampsia, hypoglycemia in pregnancy, hyperlipidemia or diabetes, pregnancy weight gain, weight of the previous offspring, and gender of the newborn were evaluated. Besides, family history of disease was also recorded.

Restricting systolic and diastolic arterial blood pressure (SABP and DABP, respectively) was measured in all the mothers after they had been sitting for at least 30 min. A mercury sphygmomanometer was used to measure blood pressure thrice at the right arm. Eleven mothers were diagnosed with hypertension (blood pressure equal to or greater than 140/90 mm Hg) during pregnancy, without clinical evidence of edema or proteinuria, after 20 weeks of gestation.

Birth outcomes. Infants were classified as SGA, LGA, or AGA according to the following criteria: SGA was defined as birth weight for gestational age less than the 10th centile of a published Argentinian fetal growth reference, and LGA was defined as birth weight above the 90th centile for gestational age and sex.14 The rest of the newborns were considered as AGA.

Evaluation of the newborn also included physical examination, search for any features suggestive of a malformation syndrome, supine lengths, Apgar score evaluation, and type of delivery.

Biochemical Measurements. Blood was drawn from pregnant women who were in a supine resting position for at least 30 min and from the umbilical cord at delivery. Routine laboratory variables were measured using standard clinical laboratory techniques by the time of the delivery. A commercial ELISA kit was used to measure plasma leptin concentrations (Assay Designs, Inc. U.S.A.) in blood samples collected with sodium EDTA; concentrations of plasma total homocysteine were determined using immunochemoluminsence methods according to the manufacturer’s instructions (Immulite 1000, DPC, Los Angeles, CA, USA).

DNA isolation and measurements of telomere length. Genomic DNA was extracted from white blood cells from a blood sample of the mothers and their babies using a previously described standard method41. An assay based on real-time quantitative PCR (iCycler thermocycler; Bio-Rad, Hercules, CA) was used for relative TLT measurement based on a method that uses a ratio of telomere DNA signal intensity to human β-globin single-copy gene signal intensity35; SYBR Green was used as a fluorescent dye (Invitrogen, Buenos Aires, Argentina).

All samples for both the telomere and single-copy gene (HBB, human β-globin) reactions were done in triplicate in the same plate, and a control DNA sample was added to each plate to account for interassay variability. The Ct value corresponds to the cycle number at which the fluorescence generated within a reaction well exceeds a predefined threshold value (100 in each run). We used the software LinRegPCR45 (version 2012.0) for efficiency computation and computed the Eff of each replicate. We determined the mean value of the replicates for each sample and each gene. The relative T/S ratio is the ratio of the telomere to HBB in one sample, set in relation to the T/S ratio in the control DNA. The formula for calculation of the relative T/S ratios can be stated as follows: T/S ratio = [Eff telomere (control)/Eff telomere (sample)]/[Eff HBB (control)/Eff HBB (sample)]. Specificity of amplification and absence of primer dimers were confirmed by melting curve analysis at the end of each run. Primer sequences and resulting PCR product lengths are shown in Table 3.

The PCR efficiencies for telomere and HBB amplification were 1.76 and 1.95 on average. As individual samples were analyzed in triplicate and accepted only if the standard deviation of the Ct values was <1 Ct, the inter- and intra- experimental coefficient of variation of the qPCR was less than 8% and 3%, respectively.

Polymorphism genotyping. Two SNPs (rs9419958 at 10:105675945 in OBF1C and rs3027234 at 17:8136092 in CTIC1) were analyzed in the naive sample population. Genotypes of these genetic polymorphisms were examined using PCR-based restriction fragment length polymorphism (RFLP); primers used in PCR-RFLP for amplifying genomic DNA are shown in Table 4. The restriction digestion of rs9419958 and rs3027234 and PCR products was carried out with restriction enzymes HpaII and Ddel (New England Biolabs Inc.), respectively, at 37°C for about 12 h. The digested products were loaded and visualized on 2.5% agarose gel after staining with ethidium bromide. Genotyping accuracy was assessed using inclusion of duplicates and negative.

The integrative analysis of biological networks was performed by the VisANT 4.0 platform (a freely available and Web-based workbench at http://visant.bu.edu)47.

Statistical Analysis. Otherwise indicated, quantitative data were expressed as mean ± SE, and differences between groups were assessed using ANOVA or ANCOVA. Because some variables, particularly the LTT/HBB ratio, do not have a normal distribution, we assessed differences in this variable using Mann-Whitney nonparametric test, or post hoc Newman-Keuls after ANOVA or ANCOVA, after a Log transformation if applicable. Correlation between two variables was performed using Spearman rank correlation or Pearson tests. To perform these analyses, we used the CSS/Statistica package program, StatSoft V 6.0 (Tulsa, USA).

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**Author contributions**

M.T. and T.F.G.: preformed molecular experiments in human samples, J.A. and C.G.: collected clinical information, C.J.P.: conceived and designed the study, supervised the experiments, analyzed the data, wrote the MS.

**Additional information**

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