Fetal cell microchimerism: a protective role in autoimmune thyroid diseases

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Abstract

Objective: The physiological persistence of fetal cells in the circulation and tissue of a previously pregnant woman is called fetal cell microchimerism (FCM). It has been hypothesized to play a role in systemic autoimmune disease; however, only limited data are available regarding its role in autoimmune thyroid disease (AITD).

Design: Circulating FCM was analyzed in a large series of previously pregnant women with Graves’ disease (GD), Hashimoto’s thyroiditis (HT), or no disease (healthy controls (HCs)). To exclude the possible bias related to placental factors, the polymorphic pattern of human leukocyte antigen-G (HLA-G) gene, which is known to be involved in the tolerance of fetal cells by the maternal immune system, was investigated.

Methods: FCM was evaluated by PCR in the peripheral blood, and the Y chromosome was identified by fluorescence in situ hybridization in some GD tissues. HLA-G polymorphism typing was assessed by real-time PCR.

Results: FCM was significantly more frequent in HC (63.6%) than in GD (33.3%) or HT (27.8%) women ($P = 0.0004$ and $P = 0.001$ respectively). A quantitative analysis confirmed that circulating male DNA was more abundant in HC than it was in GD or HT. Microchimeric cells were documented in vessels and in thyroid follicles. In neither GD/HT patients nor HC women was the HLA-G typing different between FCM-positive and FCM-negative cases.

Conclusion: The higher prevalence of FCM in HC as compared to GD and HT patients suggests that it plays a possible protective role in autoimmune thyroid disorders. Placental factors have been excluded as determinants of the differences found. The vascular and tissue localization of microchimeric cells further highlights the ability of those cells to migrate to damaged tissues.

Introduction

During pregnancy, a bidirectional exchange of cells has been observed between the fetus and the mother, which starts between the 4th and 6th weeks of gestation (1). The passage of cells from the fetus to the mother is called fetal cell microchimerism (FCM), whereas that occurring from the mother to the fetus is named maternal cell microchimerism (MCM). Both fetal and maternal microchimeric cells have been shown to persist in the maternal circulation or tissue for decades and to have progenitor-like properties, because they are able to differentiate into different cell types (2, 3, 4). The most widely used procedure for evaluating FCM is assessing by PCR the
presence of male cells in a woman with a previous male pregnancy. By this and other methods, FCM has been extensively studied in autoimmune diseases, particularly systemic ones, because they are more often observed in females and appear to be modulated by pregnancy (5, 6, 7, 8). As far as autoimmune thyroid disease (AITD) is concerned, FCM has been documented in Hashimoto’s thyroiditis (HT) and Graves’ disease (GD) (9, 10, 11, 12, 13). These diseases occur more frequently in women, with a peak incidence during the fertile age. They appear to be related to pregnancy and often have a spontaneous resolution after delivery and an onset or exacerbation in the postpartum period, although their relation with parity is still controversial (14, 15, 16). In the few available series, which include a limited number of cases, FCM was found to be represented more at the tissue level in HT and GD as compared to non-AITDs (4, 9, 10, 11, 12, 13, 17). Even at the peripheral level, scanty and discrepant results have been reported. In particular, one study that used a PCR-based technique showed that the number of circulating fetal microchimeric cells did not differ between GD and controls (12), whereas another study that used a fluorescence in situ hybridization (FISH) analysis demonstrated that a higher number of circulating fetal cells were found in GD and HT patients as compared to healthy volunteers (18).

Thus, in the present study, we compared the prevalence of FCM in two large populations of parous women with a previous male pregnancy, one including women affected with GD or HT and the other including healthy controls (HCs). Moreover, in order to investigate if possible differences in the prevalence could be the result of placental factors, we assessed the polymorphic pattern of the human leukocyte antigen-G (HLA-G) gene. Indeed, HLA-G molecules are non-classical HLA class I antigens that are expressed by villous trophoblasts and are involved in the reprogramming of the local maternal immune response because they inhibit the activation of decidual T and NK cells; they have been proposed to positively influence the outcome of pregnancy (19, 20). The HLA-G expression profile can be modulated by genetic polymorphisms, mainly two variants at the 3’ UTR, the 14 bp insertion/deletion (rs1704), and the +3142C>G (rs1063320), which influence mRNA stability and protein production and thereby lead to increased/reduced HLA-G expression and soluble HLA-G protein amounts in body fluids (21, 22, 23). In particular, the 14 bp deletion in hetero- or homozygosis is associated with a higher HLA-G expression (high secretor profile), whereas both the 14 bp insertion and the +3142C>G polymorphism in exon 8 decrease HLA-G expression (low secretor profile). We speculated that a high secretor HLA-G genetic profile, which allows for a stronger control of maternal immune response, might favor the passage of fetal cells to the mother during pregnancy and result in a higher prevalence of FCM. Thus, the HLA-G profile has been correlated with the microchimeric status in GD, HT, and HC.

Subjects and methods

Patients

Peripheral blood was collected from 105 female patients with AITDs treated at our institution. To be enrolled, patients must have had at least one male pregnancy preceding the diagnosis of AITD. In particular, 69 women had GD, with a mean age at enrollment of 53.3 years (range 27–85), and 36 had HT, with a mean age at enrollment of 53.1 years (range 33–88). Graves’ orbitopathy (GO) was diagnosed in 58/69 GD and in 4/36 HT patients. Peripheral blood was also collected from 66 HCs with one or more male children without neoplastic or autoimmune diseases (mean age 48 years, range 29–80). Moreover, none of the control women had a thyroid disease, as assessed by FT4, TSH, anti-thyroglobulin and anti-thyroidperoxidase autoantibodies, and neck ultrasound. None of the included subjects had a history of other potential sources of chimerism, such as blood transfusion, organ transplant, twin siblings, or abortion. All of the patients gave informed consent to enter the trial, which was approved by the local ethical committee.

DNA extraction from whole-blood and formalin-fixed paraffin-embedded tissues

Genomic DNA was extracted from blood samples using an Illustra DNA Extraction Kit BACC2 (GE Healthcare, Buckinghamshire, UK). DNA was also extracted from the formalin-fixed paraffin-embedded thyroid tissues of two women with GD submitted to thyroidectomy by means of a commercial kit (Puregene Core Kit A, Qiagen Sciences).

PCR for SRY sequence

For the detection of the human Y chromosome, DNA was submitted to two rounds of PCR (35 cycles each) using primers specific for the SRY locus and corresponding to the region upstream of the SRY-coding region. We previously demonstrated that this method has a sensitivity of 1 male cell/1 million female cells (24). Each PCR analysis was
repeated twice, two negative controls (DNA from two prepubertal girls) and two positive controls (DNA from two men) were included, and special care was taken to avoid external contamination. In particular, all of the samples were handled by a female technician, positive displacement micropipettes were used, and DNA extraction, PCR preparation, and analyses were conducted in separate rooms.

**Male DNA quantification**

DNA pools were obtained by mixing the same amount of DNA extracted from ten women with HT, ten women with GD, and ten HC who were previously found to be positive for the presence of FCM by PCR amplification. Each DNA pool was precipitated and concentrated to 1 μg/μl, and sequential dilutions were conducted from 6600 ng (1 000 000 gEq) to 0.0066 ng (1 gEq). Male DNA quantities were expressed in DNA genome equivalents (gEq) by using a conversion factor of 6.6 pg DNA per cell (1 gEq = DNA content of a single cell). Dilutions obtained for each pool were subjected to two rounds of PCR. Triplicate analyses were performed for dilutions ranging from 66 to 0.0066 ng in order to improve accuracy and test reliability. Samples were electrophoresed in 3% agarose gels and visualized in situ to improve accuracy and test reliability. Samples were performed for dilutions ranging from 66 to 0.0066 ng (1 gEq). Male DNA quantities were expressed in DNA genome equivalents (gEq) by using a conversion factor of 6.6 pg DNA per cell (1 gEq = DNA content of a single cell). Dilutions obtained for each pool were subjected to two rounds of PCR. Triplicate analyses were performed for dilutions ranging from 66 to 0.0066 ng in order to improve accuracy and test reliability. Samples were electrophoresed in 3% agarose gels and visualized and photographed on an u.v. trans-illuminator (VisiDoc-It Imaging System, UVP, Cambridge, UK).

**Fluorescence in situ hybridization**

The presence of microchimeric cells was also tested at the tissue level in two GD women who were analyzed for FCM by PCR amplification. Each DNA pool was precipitated and concentrated to 1 μg/μl, and sequential dilutions were conducted from 6600 ng (1 000 000 gEq) to 0.0066 ng (1 gEq). Male DNA quantities were expressed in DNA genome equivalents (gEq) by using a conversion factor of 6.6 pg DNA per cell (1 gEq = DNA content of a single cell). Dilutions obtained for each pool were subjected to two rounds of PCR. Triplicate analyses were performed for dilutions ranging from 66 to 0.0066 ng in order to improve accuracy and test reliability. Samples were electrophoresed in 3% agarose gels and visualized and photographed on an u.v. trans-illuminator (VisiDoc-It Imaging System, UVP, Cambridge, UK).

Image analysis was performed using Leica CW4000-FISH software version Y1.3.1.

**HLA-G polymorphism typing**

In 96/105 AITD cases and in 58/66 controls, the *HLA-G* 14 bp (14 bp Ins/Del) and the +3142C>G polymorphisms were genotyped by real-time PCR, as previously described (26, 27). Briefly, 100 ng of genomic DNA were amplified in a 25 μl reaction, and the analysis performed using a 7300 Real-Time PCR System (Applied Biosystems). To analyze the 14 bp polymorphism, the forward primer 5’-GTGATGGGCTGTTAAAAGTGCACCC-3’ and the reverse primer 5’-GGAAGGAAATGCAGTTCACGATGA-3’ were used. The probe used for detection of the 14-bp deletion allele was 5’-VIC-GAGTGCAAGTTCCTTGTG-BHQ-3-3’, and the probe for the 14-bp insertion allele was 5’-FAM-CAAGATTTGTCATGCTTCC-BHQ-1-3’. To genotype samples for the +3142C>G polymorphism, the forward primer 5’-CTTTTAATTACCCCATACCTCCTCTT-3’ and the reverse primer 5’-TGTCGGCTC-TCTGCTCAAATTT-3’ were used. The MGB probe used for the detection of the 3142C allele was 5’-VIC-TAGTTTAGCTCAGTGGAC-3’, and the MGB probe for the 3142G allele was 5’-FAM-TAGTTTATHCTCAGTGGAC-3’. Each probe had a non-fluorescent quencher at the 3’ end. Amplification was performed with 0.625 μl Assay Mix 40× (Applied Biosystems) and 12.5 μl PCR master mix 2×. Before the amplification, a pre-read run was performed for 1 min at 60 °C. The amplification protocol was: an initial step for 10 min at 95 °C followed by amplification for 15 s at 92 °C and for 60 s at 60 °C for 40 cycles. A post-run reading was performed for 1 min at 60 °C.

**Statistical analysis**

Possible differences in the clinical features (age at diagnosis/enrollment, age at the first male child birth, number of sons, and period between the birth of the first male child and diagnosis/enrollment) between GD or HT patients and HC were assessed by t-test and χ² test, as appropriate. The differences in the prevalence of FCM between women with GD, HT, or HC and the association of FCM with either clinical features or the *HLA-G* profile were tested by Fisher or χ² tests. Odds ratios (ORs) and 95% CI were used to assess the strength of the association between the presence/absence of FCM and the presence/absence of the disease. Differences between values were considered significant when P<0.05. All of the tests were performed using MedCalc Software version 13.2.2 (B-8400, Ostend, Belgium).
Results

Clinical features and blood FCM in patients with GD, patients with HT, and controls

Neither the GD nor the HT group significantly differed from the HC group with respect to age at the first male birth, the period between the birth of the first male child and the diagnosis of AITD, the age at diagnosis or enrollment, and the number of male children (data not shown). The presence of male DNA of presumed fetal origin had a significantly higher prevalence in HC cases than it was in either GD cases (63.6 vs 33.3%, $P=0.0004$, OR 3.5, 95% CI 1.723–7.11), HT cases (63.6 vs 27.8%, $P=0.0005$, OR 4.55, 95% CI 1.877–11.03), or AITD women as a whole (63.6 vs 31.4%, $P<0.0001$, OR 3.818, 95% CI 1.995–7.308) (Fig. 1A and B). Among either AITD patients or HC, the cases that were positive for the presence of FCM (FMC +ve) were not significantly different from those without FCM (FMC –ve) in terms of age at the birth of first male child, number of male children, interval between the birth of the first male child and diagnosis or enrollment, age at diagnosis or enrollment, and, for patients, the interval between the diagnosis and the enrollment and the association with orbitopathy (Supplementary Table 1, see section on supplementary data given at the end of this article). In GD cases, the type of treatment (anti-thyroid drugs, thyroidectomy, or radiometabolic treatment) and the interval between the thyroidectomy or radioactive iodine ablation and enrollment were also considered, because the ablation of the thyroid tissue could have determined the disappearance of circulating fetal cells, but no differences were noted between FMC +ve and FMC –ve cases (data not shown).

Interestingly, the SRY amplification was successful during the first round PCR in the majority of HCs, but only after a two-round PCR in GD and HT cases, which indicates that there is a higher amount of circulating male DNA in controls than there is in AITD patients. Thus, with the aim of performing a quantitative evaluation, FCM +ve DNA of ten HT patients, ten GD patients, and ten HCs were pooled, and sequential dilutions were done. Interestingly, this quantitative analysis confirmed that male DNA was more abundant in HCs than it was in either GD or HT patients. In particular, in HCs, male DNA could be clearly detected up to a dilution that corresponded to 66 ng total DNA (10 000 genome equivalents), whereas it was hardly or not detected at that dilution in GD and HT cases (Fig. 2).

Identification of FCM in the thyroid tissue of women affected with GD

The thyroid tissue of two GD women, one of whom was positive and one of whom was negative for FCM at the peripheral level, was studied. In both cases, male cells of presumed fetal origin were documented in the tissue by both PCR and FISH analysis. Interestingly, microchimeric cells were detected as forming part of thyroid follicles, interposed between female maternal cells (Fig. 3A, B and C).
Moreover, a microchimeric male cell was identified in a blood vessel (Fig. 3D).

**HLA-G genetic polymorphism typing**

The *HLA-G* genetic polymorphism typing was not different between FCM +ve and −ve patients of the GD, HT, or HC groups or in the AITD cases considered as a whole. In particular, no differences were found in the prevalence of the homozygous genotype (Del/Del) and the Del allele, which correspond to a high secretor profile (Table 1). As far as the +3142C>G variant was concerned, no differences were observed between the HT or GD patients and the HCs for both the genotype distribution and the allele frequencies (data not shown).

**Discussion**

In the present large series of AITD cases, the prevalence of FCM was found to be significantly higher in HCs than it was in AITD patients, which suggests a protective role for FCM toward the onset of these diseases. Most of the previous studies in systemic and non-systemic autoimmune diseases were done at the peripheral blood level, and contrasting data have been obtained about the possible role of FCM. Some authors considered it to be pathogenic because they found a higher prevalence of FCM in affected women as compared to controls (5, 6, 7, 8) and because of the hypothesis that microchimeric cells might elicit an intra-organ graft vs host rejection (GvHR) (5, 9, 28, 29). Other authors did not confirm these data (30, 31, 32, 33) and did not support the causative role for this phenomenon, which is also observed in non-autoimmune diseases (17, 24, 25, 34, 35, 36, 37, 38, 39).

The protective role of FCM in AITD that was suggested by the present data is consistent with what has been reported for neoplastic diseases, and in particular for thyroid and breast cancers (24, 25, 35, 37). Based on the immunophenotypic characterization of FMC in neoplastic diseases, we and others hypothesized that fetal cells could migrate from bone marrow niches through blood vessels to reach the injured areas and to differentiate into cells that are able to repair the diseased tissues because at least some of the engrafted fetal cells have stem cell potential (4, 25). Moreover, further evidence to support this ‘protective’ hypothesis has come from several studies in animal models. In particular, the selective homing of fetal cells to the site of organ injury and not to healthy tissues suggests that fetal cells sense specific signals, which enables them...
to target diseased tissue (40, 41, 42). Consistently in the GD cases in the present study, male cells were documented to ‘travel’ into a blood vessel and were found interposed between female cells and forming follicles in the thyroid. Although we cannot exclude a role for fetal microchimeric cells as being innocent bystanders that do not have any significant biological effect at the thyroid tissue level in GD, two tentative hypotheses can be drawn. First, male-activated fetal T cells, monocytes, macrophages, and NK cells could be the effectors of autoimmunity and could be involved in the initiation of autoimmune diseases (allo-autoimmunity), but this hypothesis seems to be unlikely because we have not observed male cells within lymphocytic infiltration in the affected thyroid gland. Alternatively, these cells may be recognized as being partially alloimmune, that is, as giving rise to an immune reaction (auto-alloimmunity) in which the effectors are maternal immune cells. Nevertheless, the number of male cells observed in the thyroid glands of these women was too low to support this hypothesis. Second, male microchimeric cells could be involved in the repair/repopulation process because of their follicular localization and their morphology, which resembles that of thyrocytes. Indeed, the presence of male cells interposed between female cells in thyroid follicles, which has already been observed in thyroid cancer (25), in nodular goiter specimens (11), and in the normal thyroids of healthy women (17), seems to reinforce the findings at the blood level and to suggest a protective/repairing role for FCM. Nevertheless, it should be noted that the microchimeric status at the peripheral level may or may not reflect that of the thyroid tissue, as was already observed in thyroid cancer (24). Consistently, Y chromosome cells were found in both of the GD tissues that were analyzed, even though only one of those patients was positive for FCM at the peripheral level. The possibility of extensively studying microchimeric status at the tissue level in AITD is strongly limited by the fact that thyroidectomy is not indicated for the treatment of HT, and it is seldom used for GD patients, who are instead mainly treated with radioiodine.

It should be noted that contradictory data to those presented here were previously obtained by Lepez et al. (18) in a more limited AITD series. Four hypotheses can be drawn to tentatively explain these contrasting results. The first involves the different intervals between diagnosis and enrollment, which was a maximum of 5 years in the Lepez et al. study and ranged from a few months to 30 years in the present study. This longer latency could theoretically explain the differences in the amount of circulating male cells, which could be progressively eliminated over time by the autoimmune process. However, we did not find any correlation between the presence/absence of FCM and the length of the interval between diagnosis and enrollment. The second hypothesis involves the interval between the birth of the youngest son and the age at diagnosis, which was 3–12 months in the Lepez et al. study and 1–60 years in the present study. This also could have contributed to differences in the amount of circulating male cells, which may progressively lessen after delivery. Nevertheless, we did not find any correlation between the presence/absence of FCM and the length of this interval either in previous studies (24) or in the present study. Third, Lepez et al. used FISH analysis, which allowed them to identify intact male cells, whereas our PCR-based method may have also led to the detection of free-DNA released from damaged cells. Still, our qualitative results were also nicely confirmed in the quantitative analysis, which demonstrated a higher amount of male DNA in HCs, and we estimate that our

| Table 1 HLA-G genetic polymorphism typing in healthy controls (HC), Hashimoto’s thyroiditis (HT) patients, and Graves’ disease (GD) patients as either positive or negative for fetal cell microchimerism (FCM). |
|---|---|---|---|
| **14 bp insertion/deletion polymorphism** | **GD n (%)** | **HT n (%)** | **HC n (%)** |
| | FCM +ve | FCM −ve | P | FCM +ve | FCM −ve | P | FCM +ve | FCM −ve | P |
| **Genotype** |  |  | 0.41 |  | 0.12 | 0.97 |  | 0.57 | 0.10 | 0.9 |
| Ins/Ins | 4 (18.1) | 9 (23.1) |  | 2 (20) | 1 (4) |  | 7 (19.4) | 4 (18.2) |  |
| Del/Del | 6 (27.3) | 10 (25.6) |  | 2 (20) | 13 (52) |  | 7 (19.4) | 5 (22.7) |  |
| Ins/Del | 12 (54.5) | 20 (51.3) |  | 6 (60) | 11 (44) |  | 22 (61.1) | 13 (59.1) |  |
| Total | 22 | 39 |  | 10 | 25 |  | 36 | 22 |  |
| **Allele** |  |  | 0.57 |  | 0.10 | 0.9 |
| Ins | 20 (45.5) | 40 (51.3) |  | 10 (50) | 13 (26) |  | 36 (50) | 21 (47.7) |  |
| Del | 24 (54.5) | 38 (48.7) |  | 10 (50) | 37 (74) |  | 36 (50) | 23 (52.3) |  |
| Total | 44 | 78 |  | 20 | 50 |  | 72 | 44 |  |

+ve, positive; −ve, negative.
analysis was derived mostly from intact cells. Finally, the discordancy could be the result of the low number of patients enrolled by Lepez et al. (11 women) or to the non-selection of control cases. On the contrary, control cases were well selected in the present study by biochemical and ultrasound examinations in order to exclude non-clinically evident thyroid diseases.

In the present study, the 3' UTR polymorphisms 14 bp insertion/deletion (rs1704) and +3142C>G SNP (rs1063320) that control HLA-G expression (21, 22) were studied for the first time inAITD. Indeed, because high levels of HLA-G expression seem to be crucial for successful implantation, we hypothesized that a high secretor genetic profile in the mother could be associated with a higher tolerance of fetal cells and is thus associated with higher levels of FMC. No differences in the HLA-G typing or, in particular, in the secretor genetic profile, were found between FCM +ve and −ve cases in the GD, HT or HC groups, which is consistent with data previously obtained in HTs and controls (43). Thus, the present findings allowed us to exclude the suggestion that the higher prevalence of FCM in HCs could be the result of a high secretor genetic profile that leads to a facilitated passage of fetal cells. Along these lines, a recent study tested the same hypothesis by analyzing HLA-G polymorphisms in children from women with scleroderma (44) and showed that a high secretor genetic profile does not allow more fetal-to-mother chimerism traffic.

In conclusion, circulating fetal cells are significantly more abundant in HCs than they are in patients with thyroid autoimmunity, which suggests a protective role for FCM inAITDs. The possible contribution to these findings of a high secretor HLA-G genetic profile has been excluded. The localization studies confirmed that microchimeric cells have the ability to migrate to diseased tissues and may have a possible protective/repairing function.

References
