Expression of a truncated form of KIT tyrosine kinase in human spermatozoa correlates with sperm DNA integrity

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BACKGROUND: TR-KIT, a truncated form of KIT (the KITL receptor), corresponding to the c-terminal half of the intracellular split tyrosine kinase domain, is expressed during the haploid stages of mouse spermatogenesis, and is one of the candidate sperm factors possibly involved in egg activation at fertilization.

METHODS: Immunocytochemistry of adult human testis, and studies of human semen samples from volunteer donors through immunofluorescence, confocal microscopy, flow cytometry, western blot and RT–PCR analyses were performed.

RESULTS: We show that the TR-KIT is expressed during spermiogenesis in the human testis, and that it is maintained in human ejaculated spermatozoa. TR-KIT is localized both in the equatorial segment and in the sub-acrosomal region of the human sperm head. The equatorial localization of the TR-KIT persists after the spontaneous acrosome reaction. Cytometric analysis of several sperm samples from volunteer donors, showed variable degrees of the TR-KIT-specific immunolabeling, and a significant inverse correlation (Pearson’s coefficient, $r = -0.76, P < 0.0001, n = 23$) of the TR-KIT positivity with markers of sperm damage, i.e. DNA fragmentation, as revealed by terminal deoxynucleotidyl transferase-mediated deoxyuridine triphosphate-nick end labeling (TUNEL) analysis and the intense clusterin positivity. We also found less significant inverse correlation with altered head morphology ($r = -0.47, P < 0.05, n = 23$) and direct correlation with sperm forward motility parameters ($r = 0.59, P < 0.01, n = 23$).

CONCLUSIONS: The TR-KIT is present in the equatorial region of human spermatozoa, which is the first sperm component entering into the oocyte cytoplasm after fusion with the egg. This localization is consistent with the function previously proposed for this protein in mice. In addition, the TR-KIT represents a potential predictive parameter of human sperm quality.

Key words: TR-KIT / human spermatozoa / equatorial segment / DNA damage / clusterin

Introduction

The KIT receptor tyrosine kinase (KITL receptor) plays an essential role in the mitotic stages of mouse spermatogenesis (Sette et al., 2000; Rossi et al., 2000, 2003; Pellegrini et al., 2008), but with the onset of meiosis its expression, at both the RNA and protein level ceases. After meiosis the TR-KIT, a truncated form of KIT corresponding to the c-terminal half of the intracellular split tyrosine kinase domain, is expressed beginning at the round spermatid stage (Sorrentino et al., 1991; Rossi et al., 1992). The mouse TR-KIT is originated by transcription of an alternative mRNA from a cryptic promoter in the 16th intron of the mouse KIT gene, which is active during spermiogenesis (Albanesi et al., 1996). TR-KIT localizes to the residual cytoplasm, midpiece and the post-acrosomal region of the mouse epididymal spermatozoon (Albanesi et al., 1996; Sette et al., 1997). Micro-injection of the recombinant TR-KIT into metaphase II-arrested mouse oocytes triggers events of early embryogenesis, such as cortical granule exocytosis, completion of the second meiotic division, formation of pronuclei, and development up to the morula stage (Sette et al., 1997). The TR-KIT-mediated parthenogenetic egg activation...
requires its interaction with Fyn (a Src-like kinase) and the adaptor protein Sam68, with the consequent activation of phospholipase Cγ1 within the egg cytoplasm (Sette et al., 1998, 2002; Paronetto et al., 2003). Interestingly, Fyn and/or other egg-carried Src-like kinases have been shown to play a critical role in egg resumption from meiosis II at fertilization and the subsequent zygotic development in mammals (Meng et al., 2006; McCinnis et al., 2007; Reut et al., 2007; Tomashov-Matar et al., 2008; Luo et al., 2009; Levi and Shalgi, 2010).

Thus, the TR-KIT has been proposed as a sperm factor possibly involved in sperm-mediated egg activation at fertilization (Rossi et al., 2003). However, it cannot be excluded that the TR-KIT, besides its eventual participation to the egg activation process at fertilization, plays additional roles during mouse spermatogenesis or in the mature sperm cell function.

Restricted expression of KIT in the membrane of spermatogonia and of TR-KIT in the cytoplasm of spermatids was also observed in the rat seminiferous epithelium, both at the RNA and protein level (Prabhu et al., 2006). As in rodents, in the adult human testis full-length KIT is expressed in spermatogonia, as revealed by the use of antibodies directed against the KIT extracellular domain (Natali et al., 1992; Strohmeyer et al., 1995; Sandlow et al., 1996). However, no clear information is available about the possible expression of TR-KIT in post-meiotic stages in human testis. Immunocytochemical analysis using a polyclonal antibody directed against the human KIT c-terminus (Santa Cruz, C-19, cat. no. sc-168), which recognizes both full-length KIT and TR-KIT, showed positivity in spermatogonia and in elongating spermatids of marmoset monkeys, but not in meiotic spermatocytes, similarly to reported observations in rodents (von Schonfeldt, 1999). Using the same antibody, Feng et al. (2005) reported immunocytochemical reactivity at the level of the acrosome in the human sperm head, but, on the basis of western blot analysis, they claimed that this positivity was due to the 150 kDa full-length KIT. In a recent study, using another antibody directed against the human KIT c-terminus (Dako, Anti-Human CD117, cat. no. A4502), KIT positivity within the human testis was confined to spermatogonia and spermatids, whereas no staining was found in meiotic spermatocytes, nor in Sertoli cells, suggesting a pattern of expression of the two alternative products of the KIT gene analogous to that observed in rodents (Unni et al., 2009). Even more recently, by using a further polyclonal antibody directed against the c-terminus of human KIT (Abcam, cat. no. ab16832), KIT immunoreactivity in human spermatids was also evident (He et al., 2010).

A truncated form of KIT analogous to mouse TR-KIT is aberrantly expressed in human cancer cell lines of various origin and in primary prostate malignant tumors both at the RNA and protein level (Toyota et al., 1994; Takaoka et al., 1997; Paronetto et al., 2004). The aberrant transcript encoding human TR-KIT originates from a cryptic promoter present within the 15th intron of the human KIT gene (Toyota et al., 1994; Paronetto et al., 2004), and not from the 16th intron, which appears to be the functional TR-KIT promoter only in mice (Sakamoto et al., 2004). Interestingly, murine hematopoietic stem cells and multipotent progenitors express the TR-KIT at the RNA and protein level, so that a potential role for this protein in the self-renewal of these cells has been postulated (Zayas et al., 2008). The TR-KIT ability to activate early embryonic development when microinjected into mouse eggs, its expression in hematopoietic stem cells, and its aberrant expression in human tumors, in which it activates Src-like kinase activity (Paronetto et al., 2004), suggest an important role of these intracellular alternative KIT gene products in regulating developmental events involving growth-factor-independent cell proliferation.

Among assisted reproduction techniques (ART), the intracytoplasmic sperm injection (ICSI) is overall considered a ‘safe’ technique, however, it bypasses the natural mechanisms of gamete selection. In particular, the injection of spermatozoa-carrying DNA damage might be dangerous in terms of potential malformations in the conceived child (Lewis, 2002). Moreover, DNA damage in the male germ line, mostly due to oxidative stress (Agarwal et al., 2008), has been associated with male infertility, failed fertilization, impaired preimplantation development and poor pregnancy outcomes, whether the insemination is natural or artificial (Irvine et al., 2000; Agarwal and Said, 2003; Aitken et al., 2009; Barratt et al., 2010). A significant increase in DNA damage can also be found in sperm from infertile men classified as normozoospermic on the basis of normal standard sperm parameters (Saleh et al., 2002). Thus, discovery of new sperm parameters predictive of DNA integrity will have an important impact on the development of ARTs.

In the present paper, we have investigated the expression of the TR-KIT in human ejaculated spermatozoa from the seminal fluids of volunteer donors. We found that TR-KIT, but not full-length KIT, is present in the equatorial and sub-acrosomal region of the human sperm head. Moreover, our results show a significant positive correlation of the TR-KIT positivity with sperm DNA integrity, and we therefore propose the TR-KIT as a new potential marker of human semen quality.

### Materials and Methods

#### Semen samples

Specimens from 23 reputedly normozoospermic volunteer donors attending the Seminology Laboratory - Sperm Bank of the Department of Medical Physiopathology, University of Rome ‘La Sapienza’, were collected as masturbation after 3–5 days of sexual abstinence, into sterile plastic jars. Material was utilized after having obtained written a consent for the utilization of semen samples. Samples were allowed to liquefy for 60 min at 37°C and were then evaluated according to WHO criteria (World Health Organization, 1999). The variables taken into consideration were: ejaculate volume (ml), sperm concentration (n × 10⁶/ml), total sperm count (n × 10⁹/ejaculate), forward motility (%) and morphology (% abnormal forms). None of the donors were found to be oligozoospermic. Of the 23 donors, 20 were within the normal standard parameters defined by WHO (normozoospermic), whereas the remaining 3 were borderline asthenoteratozoospermic.

After removing seminal plasma by centrifugation, sperm pellets were washed twice in 1× phosphate-buffered saline (PBS), counted and divided into aliquots.

For swim-up purification of semen samples, 1 μl of each seminal sample was diluted 1:2 with Hank’s medium and centrifuged for 10 min at 300 g. After centrifugation, the supernatant was discarded and an aliquot of 0.5 ml of Hank’s was layered on the pellet. The spermatozoa were allowed to migrate for 30 min at 37°C. After migration, the supernatant was gently aspirated and processed for further studies.
Immunohistochemistry and immunocytochemistry

Histological sections from two formalin-fixed paraffin-embedded human testicular normal tissue fragments obtained from orchectomy of testicular cancer patients (a gift from Dr Giovanni Bertalot, Ospedale di Lecco, Brescia, Italy) were cut into 5-μm sections and mounted on polylysine coated slides. Sections were dewaxed, rehydrated and processed in a microwave for antigen retrieval in pH 6.0 citrate buffer. After quenching of endogenous peroxidase and blocking of non-specific binding, sections were incubated for 1 h at room temperature (RT) with 1:200 polyclonal antibody directed against the human KIT c-terminus (Santa Cruz, C-19, cat. no. sc-168), which recognizes both full-length KIT and TR-KIT, extensively washed and then processed using the avidin–biotin peroxidase complex (ABC) procedure, according to the manufacturer’s protocol (UltraTek HRP Anti-Polyvalent kit, ScyTek Laboratories, USA), negative controls were performed using affinity-purified rabbit IgG or omitting the primary antibody. Peroxidase activity was revealed using 3,3-diaminobenzidine tetrahydrochloride (Roche, Italy), nuclei were counterstained with Mayer’s hematoxylin. After washing, sections were permanently mounted and observed by light microscopy.

For immunocytochemical studies on human semen samples, we used the same UltraTek HRP Anti-Polyvalent kit and protocol described above. In particular before primary antibody incubation, ejaculated spermatozoa, obtained after washing of sperm pellets and spotted into polylysine coated slides, were briefly treated with cold acetone (to fix and permeabilize cells) and air dried. Samples were mounted and observed by light microscopy.

Immunofluorescence analysis and confocal microscopy

Ejaculated spermatozoa, obtained after sperm pellet washing, were spotted onto polylysine coated slides, fixed in cold acetone and air dried. After blocking for 2 h at RT with 1× PBS/5% bovine serum albumin (BSA 5%)/normal goat serum (NGS) 10% or 10% NGS plus 10% normal donkey serum (NDS) in double-immunostaining experiments), sperm cells were incubated overnight at 4°C with 1:200 C-19 rabbit anti-human KIT antibody, extensively washed, incubated for 1 h at RT with 1:100 secondary goat anti-rabbit Alexafluor 488-conjugated antibody (Molecular Probes), washed, mounted and observed at fluorescence microscopy (Axioplan 2 Imaging system, Carl Zeiss, Germany). Negative controls were performed omitting primary antibodies and using the affinity-purified rabbit IgG. Nuclei were stained using TOTO-3 dye (1:2000; Sigma Aldrich, Italy). In order to assess the specificity immunobinding of C-19 polyclonal antibody to the human intracellular KIT sperm antigen, before the immunostaining, primary antibody was pre-adsorbed with a 100-fold mass excess of the immunogenic cognate peptide (Santa Cruz Biotechnology) and then used in paired control slides to evaluate the efficiency of competition; this experiment was repeated three times on different semen samples.

In order to perform double-immunostaining for the TR-KIT and acrosin, incubation for 1h at RT with 1:500 mouse monoclonal anti-human acrosin antibody (Biospora Corp., Chile) was performed after O/N C-19 anti KIT incubation, whereas secondary antibodies, 1:1000 secondary Alexafluor 488 goat anti-rabbit (Molecular Probes, Invitrogen) and 1:300 donkey Cy3-conjugated anti-mouse (Jackson ImmunoResearch, UK), were used together for 1h at RT. Nuclei were stained using TOTO3 dyes (1:2000; Molecular Probes, Invitrogen). Labeled-sperm cells were observed with a Leica laser scanning microscope TCS SP2 equipped with three laser lines. Each channel was acquired separately using specific laser lines to avoid a bleed-through of the fluorochromes. Photomicrographs were acquired using LAS AF Leica Confocal Software (Leica, Germany).

Immunoblot analysis

Human spermatozoa from raw semen of normozoospermic donors and from patients recruited for fertility studies were resuspended in lysis buffer (50 mM HEPES, pH 7.5, 150 mM NaCl, 15 mM MgCl2, 15 mM EGTA, 250 mM NaVO4, 10 mM b-glycerophosphate, 1% Triton X100, 0.1% SDS, supplemented with protease inhibitor cocktail) and sonicated for three cycles of 30 s on ice. The control recombinant mouse TR-KIT protein was prepared by protein extraction from transfected cell lines as described previously (Albanesi et al., 1996; Sette et al., 1997). Lysates were cleared by centrifugation at 10 000 g for 10 min. The supernatant fraction was collected, diluted in SDS-PAGE sample buffer and boiled for 5 min. Samples were separated on 10% SDS-PAGE gels and transferred to polyvinylidenefluoride Immobilon-P membranes (Millipore) using a semidy blotting apparatus (BioRad). Western analysis was carried out as previously reported (Sette et al., 2002) using the C-19 anti-human KIT (c-terminal) antibody (1:1000 dilution) O/N at 4°C. Competition was performed by preincubating the antibody with an excess of the immunogenic peptide (100 × molar excess; 1:1 mass ratio) for 12 h before dilution in the primary antibody solution for hybridization (Sette et al., 1997). Secondary anti-rabbit IgG conjugated to horseradish peroxidase (Amersham) were incubated with the membranes for 1 h at RT at a 1:10 000 dilution in PBS containing 0.1% Tween 20. Immunostained bands were detected by the chemiluminescent method (Santa Cruz Biotech.). The human semen sample in which 100% of spermatozoa lacked the acrosome was kindly provided by Dr Jan Tesarik (Center for Reproductive Medicine, European Hospital, Rome, Italy).

For the RNA interference analysis in transfected cell lines, a 500-bp inverted repeat corresponding to part of the S′ untranslated region and the beginning of the mouse TR-KIT open reading frame (ORF) (Rossi et al., 1992) was subcloned in the pDECAP vector described by Shinagawa and Ishii (2003). HEK293 cells were co-transfected with a recombinant plasmid expressing the mouse TR-KIT ORF (Albanesi et al., 1996; Sette et al., 1997) and a second plasmid expressing an Myc epitope (for control for transfection efficiency). Cells were also co-transfected either with the pDECAP empty vector or with increasing amounts of the interfering plasmid constructs. Protein extracts from transfected cells were processed for immunoblot analysis. Western blots were probed with the C-19 anti-human-KIT c-terminus antibody and an antibody directed against the Myc epitope.

RT-PCR analysis

In order to verify the presence of specific KIT mRNA molecules in sperm cells following osmotic shock, the total RNA was extracted from 2 × 10⁶ purified ejaculated spermatozoa as previously described (Muciaccia et al., 2007) using high pure RNA isolation kit (Roche) according to the manufacturer’s instructions. Purified RNA was treated for 15 min at 37°C with 2 units of Deoxyribonuclease I, Amplification Grade (Invitrogen-Life Technologies) to prevent DNA genomic contamination. An aliquot of total RNA was reverse-transcribed, using Senscript Reverse Transcriptase (Qiagen), a highly sensitive enzyme recommended for very small RNA amount (<50 ng) and oligo-dT as primer. Then, by PCR amplification, total RNA was tested for prostatine-2 gene (PRM-2, specifically expressed only in haploid germ cells), using a primer set able to discriminate between genomic DNA and sperm mRNA (Miller et al., 1994).

In order to amplify KIT mRNA and discriminate between different transcripts (full-length mRNA or truncated form), aliquots of cDNA were used in amplification reactions using the following oligonucleotide primers...
spanning four contiguous human KIT gene exons (Accession number: nG_007456): e15 (forward) GTACTAATGAGTACATGGACATG; e16 (forward) GCTCATACAGAAGAGATG; e17 (forward) TGTATT CACAGACTTGGCACGC; e18 (reverse) TGCTTTCAGGGCCAT CCACCTCAC.

Amplification conditions were: first denaturation at 94°C for 3 min, then 94°C for 30 s, 65°C for 45 s, 72°C for 40 s for 50–80 cycles, in 50 μl final volume using 2 U of Taq DNA polymerase (Roche). A control reaction tube, in which reverse transcriptase enzyme was omitted, was also included to test possible DNA genomic contamination. Aliquots of PCR product (i.e. 1/5 of total reaction volume) were separated on 2% agarose gel and visualized by ethidium bromide staining.

Cytometric analysis of human sperm cells

Presence of intracellular TR-KIT and clusterin antigens on freshly isolated sperm cells from presumably healthy donors was assessed by cytometric analysis. Aliquots of 10^6 sperm cells/tube were briefly fixed on ice using 2% paraformaldehyde, washed twice and then permeabilized using 0.1% sodium citrate/0.1% Triton X-100. After washing sperm cells were incubated for 30 min on ice with primary antibodies (1:200 in PBS/1% BSA, C-19 rabbit polyclonal antibody against the c-terminus of human KIT, #sc-168; H-330 polyclonal rabbit anti-human clusterin antibody, #sc-8354, both from Santa Cruz Biotechnology, Germany). The C-19 antibody recognizes both the full-length KIT and the TR-KIT, whereas the anti-clusterin antibody can recognize either the acrosomal or the surface isoforms of clusterin. Sperm cells from the same samples were also incubated in a control tube with the same concentration of the corresponding affinity-purified rabbit IgG (Sigma), as recommended by technical data sheets. Antibody-labeled sperm suspensions were washed twice with 1 × PBS/1% BSA, incubated with phycoerythrin-conjugated secondary antibody (Goat anti-rabbit-PE, by Jackson ImmunoResearch Laboratories, UK). In order to assess the presence of intracellular TR-KIT and to exclude its sperm surface expression, control tube reactions were also performed using aliquots of unfixed and unpermeabilized sperm cells in which the anti-KIT antibody immunolabeling failed to detect any surface specific signal (data not shown). The specificity of the sperm cells gate was verified by analysis of sperm DNA content using the DNA binding dye propidium iodine. For double detection of both the TR-KIT antigen and the sperm DNA fragmentation by the terminal deoxynucleotidyl transferase-mediated deoxyuridine triphosphate-nick end labeling (TUNEL) technique, aliquots of 1 × 10^6 spermatozoa were treated by In Situ Cell Death Detection Kit-Fluorescein (Roche, Italy), according to the manufacturer’s instructions before the TR-KIT immunolabeling. Cells were then run through a flow cytometer (Epics XL Beckman Coulter, USA) using an Argon 488 nm excitation laser and 5000 and 10 000 events were, respectively, acquired for each single and double reaction/tube. Results were expressed as the percentage of positive spermatozoa gated. In order to assess the specificity immuno-binding of C-19 polyclonal antibody to human intracellular TR-KIT sperm antigen, before cytometric analysis, primary antibody was pre-adsorbed with a 100-fold mass excess of the immunogenic cognate peptide (Santa Cruz Biotechnology) and then used in paired control tubes to evaluate the efficiency of competition binding reaction; the competition experiment was repeated four times on different semen samples.

Statistical analysis

Data from morphological and cytometric analyses of semen samples were utilized to build matrices for the calculation of r-values (Pearson’s product-moment correlation coefficients); the significance (two-tailed probability values) of r coefficients were calculated on the basis of the correlation values and the sample sizes.

Results

Immunohistochemistry in normal adult testis

Using a polyclonal antibody directed against the c-terminus of human KIT (Santa Cruz, C-19, sc-168), which recognizes both the full-length KIT and the TR-KIT, the KIT immunostaining was evident in the membrane of several spermatogonia in the basal layer and in interstitial Leydig cells, whereas Sertoli cells and spermatocytes inside the seminiferous tubules were negative (Fig. 1A–C). Clear positivity was also evident in the haploid stages of spermatogenesis. KIT immunostaining was evident in the cytoplasm of round (Fig. 1A and B) and elongating spermatids (Fig. 1C), sometimes with a distinct perinuclear distribution in the area of the developing acrosome (Fig. 1B). The presence of KIT immunostaining both in spermatogonia and in spermatids, but not in spermatocytes, is substantially similar to recent observations by other groups using two different polyclonal antibodies directed against the c-terminus of human KIT (Unni et al., 2009; He et al., 2010).

Analysis of ejaculated spermatozoa by immunocytochemistry

Preparations of freshly ejaculated spermatozoa from normozoospermic subjects were fixed after permeabilization and probed with the C-19 anti-human KIT antibody, followed by secondary antibody conjugated to peroxidase (Fig. 1D and E). Immunocytochemical analysis revealed KIT staining in the anterior region of the sperm head, with maximal intensity often observed in the equatorial segment (Fig. 1D), or isolated intense staining in the equatorial segment alone (Fig. 1E). However, not all sperm heads resulted positive to the KIT immunostaining (Fig. 1D). Notably, most of KIT-negative spermatozoa appeared to have mis-shaped heads or other evident anomalies extending to the midpiece (Fig. 1D).

Immunofluorescence and cytometric analysis of ejaculated spermatozoa

A typical low magnification picture of fixed-permeabilized sperm preparation from a normozoospermic patient probed for immunofluorescence with the anti-human KIT antibody (green-cyan signal), and counterstained with TOTO-3 dye to identify sperm nuclei (blue signal), is shown in Fig. 2A. Similar to immunocytochemistry observations, KIT positivity was evident in most sperm heads, especially in the equatorial segment, but, again, not all spermatozoa appeared to be positive. Figure 2B shows that the fluorescent signal was almost completely abolished if the primary antibody had been pre-incubated with a 100-fold mass excess of the immunogenic peptide, demonstrating specificity of KIT immunostaining in the sperm head.

In order to quantify positivity of KIT immunostaining, we performed cytometric analysis. A typical example of fluorescence-activated cell sorting (FACS) analysis with a permeabilized sperm sample from a normozoospermic subject is shown in Fig. 2C. Specificity of the shift (red line) was confirmed by its abolishment after pre-incubation of the primary antibody with the immunogenic peptide (Fig. 2C, blue line). KIT positivity in spermatozoa varied in different subjects, ranging from 10 to 93%, with a median of 54% (Table I). No shift was obtained with the same antibody if sperm samples had not
been permeabilized (data not shown), indicating that the KIT positivity is due to intracellular sperm components.

**Subcellular localization of KIT immunostaining in human sperm heads**

Typical double immunofluorescence analysis performed with anti-KIT (green) and anti-acrosin (red) antibodies, confirmed that the KIT–positivity in human sperm was localized in the anterior region of the head, with maximal intensity in the equatorial segment, behind the acrosome, or exclusively in the equatorial region (Fig. 3A; see also Supplementary Material, Fig. S2A). Notably, KIT equatorial staining persisted in acrosin-negative sperm heads, indicating that it is not localized in the acrosome vesicle nor released during spontaneous acrosome reaction. In some sperm heads, KIT and acrosin signals appeared to uniformly merge all over the anterior region of the head (yellow signal), with no evident accumulation in the equatorial segment. Some sperm heads were KIT-negative, but positive for acrosin staining, or negative for both signals. Careful morphological analysis of KIT positivity in sperm samples from several normozoospermic volunteer donors indicated that 50–80% of KIT-positive spermatozoa showed maximal or isolated staining in the equatorial segment, whereas the remaining percentage of sperm cells showed apparent merging of the KIT and acrosin signals in the anterior region of the head. When DNA counterstaining (blue signal in Fig. 3A) was omitted, it was evident that also in spermatozoa

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**Figure 1** Immunohistochemical and immunocytochemical studies of human testis and spermatozoa with an anti-KIT (c-terminal) antibody. Localization of KIT immunostaining in histological sections of normal adult testis (A–C) and in permeabilized spermatozoa from semen of normozoospermic donors (D and E) using the C-19 rabbit polyclonal antibody against the c-terminus of human KIT. Staining was revealed with a secondary antibody conjugated to peroxydase (hematoxylin counterstained). Light microscope images at lower (A and D) and higher (B, C and E) magnifications are shown. In (D and E), light microscope images were merged with phase contrast pictures. Arrows in (A–C) indicate positive staining in spermatogonia (SG), spermatids (ST) within the seminiferous tubules, and in interstitial Leydig cells (LC). Arrows in (D and E) indicate either absence of staining (negative) or staining in the equatorial and anterior head regions of spermatozoa.
showing uniform positivity all over the anterior region of the head, the KIT signal was more heavily concentrated in the equatorial segment (Supplementary Material, Fig. S1).

To ascertain whether KIT positivity in the anterior region of the head was due to co-localization with acrosin within the acrosomal matrix, human spermatozoa were analyzed by laser-scanning confocal immunofluorescence microscopy. As shown in the series of images in Fig. 3B, representing six sequential focal planes of a typical confocal analysis (supplementary Fig. S2B), proceeding from the proximal to the distal plane, the red acrosin signal was the first to appear and the last to disappear with respect to both the green (KIT) equatorial signals and the yellow (KIT-acrosin merge) signal in the anterior region of the head, which were closer to the plane of maximal blue staining (DNA). Thus, KIT positivity appears to be maximal in the sub-acrosomal region of the sperm heads, similar to the perinuclear staining observed in developing spermatids (cfr. Fig. 1B,C).

**Molecular identification of TR-KIT as the KIT gene product normally expressed in human spermatozoa**

Intracellular KIT immunostaining in the perinuclear region of haploid spermatids and in the sub-acrosomal-equatorial region of sperm heads, suggests that, like in rodents, human post-meiotic KIT positivity might be due to expression of the human equivalent of mouse TR-KIT. In order to clarify this point, we performed western blot analysis of protein extracts from freshly ejaculated human spermatozoa using the same antibody used for the morphological experiments. This antibody is directed against the c-terminal portion shared by both the full-length KIT and TR-KIT. A typical immunoblot analysis is shown in Fig. 4A. No signal was evident in the high molecular weight (MW) range of the blot, indicating that the 150 kDa full-length KIT is not expressed in normal human spermatozoa. However, two lower
MW bands, of approximately 50 and 30 kDa were observed. The size of the 30 kDa band coincided exactly with that of a recombinant mouse TR-KIT protein expressed in transfected cell lines. Specificity of these bands was demonstrated by their almost complete disappearance when the primary antibody had been pre-incubated with a 100-fold molar excess (1:1 mass ratio) of the competing immunogenic peptide, whereas other non-specific bands detected in transfected cell lines were not competed. Identical results were obtained in human sperm samples by using the cross-reactive anti-mouse-KIT-c-terminus antibody described by Albanesi et al. (1996; data not shown). The specificity of the antibody was confirmed by co-transfecting cell lines with a TR-KIT expressing plasmid and a series of plasmids designed to elicit RNA interference against TR-KIT mRNA (Supplementary Material, Fig. S3).

The 50 and 30 kDa TR-KIT bands were also observed when immunoblot analysis was performed with a human sperm sample in which 100% of spermatozoa lacked the acrosome, confirming that the TR-KIT is not localized inside the acrosomal matrix (Fig. 4B). Immunoblot analysis performed with randomly selected sperm samples from patients recruited for fertility studies, showed that the low MW KIT bands were observed with variable intensity in different subjects, in agreement with the observed variability of KIT immunostaining observed in immunocytochemical, immunofluorescence and cytometric studies in normozoospermic subjects (Fig. 4C). Notably, the relative intensity of the 50 versus the 30 kDa specific bands varied in different samples. Two specific TR-KIT immunoreactive bands of approximately the same MW were observed also in mouse elongating spermatids and in mouse epididymis spermatozoa, and we suggested that the 50 kDa band might be generated by a covalent interaction of the 30 kDa TR-KIT protein with some other protein/s present in haploid cells, but not in transfected cell lines (Albanesi et al., 1996).

Identification of TR-KIT as the KIT gene product normally expressed in human spermatozoa was confirmed also at the RNA level through semi-quantitative RT–PCR. Analysis of cDNA obtained from RNA extracted from human spermatozoa of fertile subjects shows that TR-KIT, but not KIT, mRNA is detectable in human sperm (Fig. 4D). Indeed, using as reverse primer an oligonucleotide corresponding to exon 18 (shared by both KIT and TR-KIT) and different forward primers, we observed the positive signal of the expected size (209 bp band) only when the primer annealed to the KIT exon 17 (which corresponds to the first exon of the TR-KIT-specific mRNA expressed during mouse spermiogenesis). By contrast, forward primers that should amplify RNA encoding the full-length KIT receptor (oligonucleotides corresponding to sequences from exon 15 and the beginning of exon 16), did not generate any RT–PCR product even after a higher number of PCR cycles.

Correlation of TR-KIT expression in the sperm heads with sperm DNA integrity

Since lack of the TR-KIT immunostaining was frequently associated with morphological anomalies of the sperm heads, we performed co-staining of KIT and DNA fragmentation, a well-defined marker of sperm head damage, through the TUNEL technique. We also performed co-immunofluorescence staining for KIT and clusterin. Indeed, in bulls, rams and humans, normal spermatozoa are stained by anti-clusterin antibodies, due to the presence of clusterin produced

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<th>No. of samples</th>
<th>Clusterin (M2) FACS positivity</th>
<th>Clusterin (M3) FACS positivity</th>
<th>TUNEL FACS positivity</th>
<th>% TUNEL/Clusterin double FACS positivity</th>
<th>% TR-KIT FACS positivity</th>
<th>% TR-KIT FACS positivity after swim-up purification</th>
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during the spermatogenic process within the acrosomal cap, whereas morphologically abnormal spermatozoa have an extensive surface coating of a different clusterin isoform all over the sperm length, including the tail, which is not detectable on normal spermatozoa (O’Bryan et al., 1994; Ibrahim et al., 2000, 2001; Martínez-Heredia et al., 2008, Muciaccia et al., unpublished data). We routinely observed that most of TR-KIT positive spermatozoa were TUNEL negative or faintly reactive for clusterin in the acrosomal cap, whereas most KIT negative spermatozoa, which often were positive for other sperm head proteins, such as acrosin, were heavily stained by the anti-clusterin antibody and mostly TUNEL positive (Muciaccia et al., unpublished data). We performed cytometric analysis of 23 sperm samples from volunteer donors, 20 of which were normozoospermic and 3 asthenoteratozoospermic (see Table I for average seminal and cytometric parameters). We observed that TR-KIT positive cells were highly enriched in the TUNEL-negative cell populations (Fig. 5A). Figure 5B shows that the percentage of shift with the anti-KIT antibody in FACS analysis was inversely related to the percentage of M3 shift obtained with the anti-clusterin antibody (corresponding to the abnormal sperm population with a very high immunofluorescence staining due to the extracellular coating of clusterin; Muciaccia et al., unpublished data). By contrast, the percentage of shift with the anti-KIT antibody was directly related to the clusterin M2 shift (corresponding to the normal sperm population, which show a moderate immunofluorescence staining due to the intracellular clusterin present in the acrosomal region).

A summary of cytometric data obtained in the 23 sperm samples is shown in Fig. 6A. The lowest levels of TR-KIT expression was found in the semen of two of the three donors, which resulted to be asthenoteratozoospermic. In these samples, we also found the highest levels of DNA damage, as revealed by the TUNEL analysis. Statistical analysis (Table II) revealed a significant (Pearson’s coefficient, \( r = -0.76, P < 0.0001, n = 23 \)) inverse correlation between the percentage of TR-KIT immunolabeling and double positivity for two markers of

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Figure 3 KIT immunostaining in human spermatozoa is mostly confined to the equatorial segment of the sperm heads and sub-acrosomal region. (A): representative example of double immunofluorescence analysis for KIT (green signal) and acrosin (red signal) in permeabilized spermatozoa from semen of a normozoospermic donor, using TOTO-3 dye to reveal DNA in sperm heads (blue signal) and transmission microscopy to check sperm morphology. Equatorial localization of KIT signal is evident in most sperm heads, and is present also in spermatozoa which had undergone a spontaneous acrosome reaction (revealed by lack of acrosin staining in the anterior head region). In a minority of cells, KIT and acrosin signals appear to merge in the anterior region of the head (yellow signal). Some sperm cells are KIT-negative but acrosin-positive, and some are negative for both antigens. Most spermatozoa with low or absent KIT reactivity display evident morphological anomalies. (B): representative laser-scanning confocal immunofluorescence microscopy images of semen of a normozoospermic donor probed with anti-KIT (green signal) and anti-acrosin (red signal) and counterstained with TOTO-3 dye to reveal DNA in sperm heads (blue signal), overlapped on the background of corresponding transmission microscopy images. Panels from 1 to 6 indicate six sequential focal planes of a typical confocal analysis. An arrow points to a sperm head in which KIT and acrosin signals appeared to merge in the anterior region of the head (yellow-orange signal) in ordinary immunofluorescence observation (supplementary Fig. S2A). In this sperm head, the red (acrosin) signal is mostly evident in the more proximal (1.2) and distal (5.6) planes, whereas the yellow-green (KIT) signals are mostly evident in the central planes (3 and 4).
Human spermatozoa express TR-KIT, but not full-length KIT. (A): representative immunoblot analysis using the C-19 rabbit polyclonal antibody against the c-terminus of the KIT or the same antibody pre-incubated with an excess of the immunogenic peptides. Protein extracts tested were (left to right) from mock-transfected HEK293 cells, from the same cells transfected with a vector expressing recombinant mouse TR-KIT, and from spermatozoa of two normozoospermic donors. High MW bands corresponding full-length KIT are absent, whereas two specific 50 and 30 kDa TR-KIT bands are evident in both sperm samples. These bands, as well the recombinant 30 kDa TR-KIT band, are almost absent after pre-incubation of C-19 with its cognate epitope. (B): immunoblot analysis of protein extracts from semen of a patient in which 100% of spermatozoa lacked the acrosome. The specific TR-KIT bands are evident also in this sample, and high MW bands corresponding full-length KIT are absent. (C): immunoblot analysis using the C-19 antibody performed with eight randomly selected sperm samples from patients (indicated by numbers) recruited for fertility studies. The 50 and 30 kDa TR-KIT bands were observed with variable intensity in different subjects and, again, high MW bands corresponding full-length KIT are absent. Equivalent protein loading was checked by reversible Ponceau staining before probing blots with the C-19 antibody. (D): a representative RT–PCR analysis of RNA extracted from spermatozoa of a normozoospermic donor, indicates that RNA encoding for TR-KIT, but not for full-length KIT, is detectable in human spermatozoa as a remnant of transcription occurring during spermiogenesis. The low MW bands present on the bottom of the lanes are due to the oligonucleotide primers used for amplification. No signals were obtained when omitting RT before the amplification step of the reaction. This experiment was repeated on RNA extracted from sperm samples from three different normozoospermic donors with similar results.
Figure 5 TR-KIT expressing spermatozoa have a low level of DNA damage and an high percentage of TR-KIT positivity corresponds to low levels of staining with anti-clusterin antibodies. (A): representative double cytometric analysis for TR-KIT immuno-positivity and DNA fragmentation on a sperm sample from a normozoospermic donor. Permeabilized spermatozoa were treated by In Situ Cell Death Detection Kit-Fluorescein (Roche, Italy) before TR-KIT immunolabeling, with the C-19 antibody. Numbers within different gate quadrants refer to the percentage of: TR-KIT positive/TUNEL negative (top left), TR-KIT negative/TUNEL negative (bottom left); TR-KIT positive/TUNEL positive (top right), and TR-KIT negative/TUNEL positive cells (bottom right). TR-KIT positive cells appear to be highly enriched in the TUNEL-negative cell population. (B): representative contemporary cytometric analysis with anti-clusterin (left panels) and anti-KIT (right panels) antibodies of permeabilized sperm samples from two different donors, showing either high (top panels) or low (bottom panels) percentage of TR-KIT positivity (M1 peaks). In the sperm sample with high TR-KIT positivity (top panels), two separate peaks of clusterin positivity (M2 and M3) are evident; the M2 clusterin peak corresponds to spermatozoa with moderate anti-clusterin staining in the acrosomal cap, whereas the M3 peak corresponds to spermatozoa with intense surface anti-clusterin staining (Muciaccia et al., unpublished data). In the sperm sample with low percentage of TR-KIT positivity (bottom panels), only the clusterin M3 peak is evident.
sperm damage, i.e. TUNEL positivity and intense clusterin immunolabeling (M3 peak). Conversely, these two negative parameters appeared to be in strong direct correlation to each other ($r = 0.96$, $P < 0.00001$, $n = 23$). In the semen samples from the same subjects, we also found a less significant direct correlation between TR-KIT positivity and sperm forward motility parameters ($r = 0.59$, $P < 0.01$, $n = 23$), and inverse correlation to the microscopically observed percentage of sperm head atypical forms ($r = -0.47$, $P < 0.05$, $n = 23$). A significant inverse correlation was also found between percentages of TR-KIT immunolabeling with increasing donors’ age ($r = -0.73$, $P < 0.0001$, $n = 23$), whereas no correlation was found between TR-KIT positivity and other semen parameters, such as sperm concentration ($r = 0.00$) or volume of the ejaculate ($r = -0.05$).

The most interesting data emerging from the FACS analysis is that the average percentage of TUNEL positivity in unselected sperm samples was $25.03 \pm 20.19$ (median value: $18.18$; $n = 23$; Table I), whereas double positivity for both TUNEL and TR-KIT dropped to $7.69 \pm 3.12$ (median value: $6.81$; $n = 17$). Thus, percentages of TR-KIT immunolabeling of human spermatozoa correlates to sperm DNA integrity, suggesting that it might constitute a positive marker of semen quality in human fertility studies. Further support to this hypothesis was obtained by comparing cytometric analysis of three total unselected sperm samples with that of the same samples after swim-up purification (see a representative example in Fig. 6B). An enrichment of TR-KIT immunolabeling was constantly observed after swim-up selections in all sperm samples tested, demonstrating that motile spermatozoa are an homogeneous population of TR-KIT immunoreactive cells (percentage of TR-KIT positivity in the three semen samples: $88.03 \pm 0.85$ in the raw semen, and $96.37 \pm 1.18$ after swim-up selection).

### Discussion

Infertile men are often characterized by a reduced sperm number, impaired sperm morphology, low sperm motility or, in many cases,
by a combination of all these factors. A less frequent cause of unexplained male infertility is failure in egg activation after sperm–oocyte fusion and/or block in the early embryonic development. The non-genomic paternal contribution to egg activation consists of triggering meiotic resumption of the oocytes from their block in metaphase II (Barroso et al., 2009). Mouse TR-KIT is one of the candidate sperm factors acting in egg activation at fertilization (Sette et al., 1997, 1998, 2002; Paronetto et al., 2003).

In this paper, we show that TR-KIT protein is expressed during human spermiogenesis and maintained in human spermatozoa, suggesting a conserved functional role for this alternative product of the KIT gene during male gametogenesis and/or at fertilization. In contrast with previous claims (Feng et al., 2005), in human spermatozoa, we found no trace of full-length KIT. This was confirmed by our finding that RNA encoding for TR-KIT, but not for full-length KIT, is present in human spermatozoa, likely as a remnant of transcription occurring during human spermiogenesis (Miller et al., 1994).

Besides TR-KIT, other candidate sperm factors for egg activation are phospholipase C zeta (PLCZETA) (Saunders et al., 2002) and post-acrosomal sheath WW domain-binding protein (PAWP) (Wu et al., 2007a). PLCZETA was shown to induce oocyte activation from the mouse sperm perinuclear matrix (Fujimoto et al., 2004). Moreover, PLCZETA is localized in the equatorial segment of bull sperm (Yoon and Fissore, 2007), and in the acrosomal, equatorial and post-acrosomal segments of human sperm (Grasa et al., 2008). PAWP exclusively resides in the post-acrosomal sheath of guinea pig and bull sperm (Wu et al., 2007a, b). Sperm deficient in oocyte activation have been shown to lack equatorial localization of PLCZETA, and immunoblot analysis showed reduced amounts of PLCZETA in sperm from infertile men (Yoon et al., 2008; Heytens et al., 2009). However, it cannot be excluded that other factors present in the perinuclear theca (PT) and/or in the equatorial region might equally be absent in spermatozoa that are unable to trigger egg activation.

PT (the cytoskeletal coat of the mammalian sperm nucleus that is removed from the sperm head at fertilization) plays a role in joining the acrosome and the post-acrosomal plasma membrane to the nucleus during spermiogenesis. In addition, it has been proposed that PT might harbor the sperm borne oocyte-activating factors (SOAFs), which trigger the signaling cascade of oocyte activation (Kimura et al., 1998; Sutovsky and Schatten, 2000; Ito et al., 2009).

The appearance and localization of the sub-acrosomal PT during human spermiogenesis is tightly associated with acrosomal biogenesis (Alvarez Sedó et al., 2009). We found that, during human spermiogenesis, the TR-KIT is strongly concentrated in the perinuclear region of spermatids and in the area of the developing acrosome. In mature spermatozoa the TR-KIT is present in the sub-acrosomal layer of the human sperm head, where the PT is found, and it is highly concentrated in the equatorial segment, similarly to PLCZETA.

Since the equatorial segment and the post-acrosomal regions are the first part of the head which fuses with the egg membrane (Florman and Ducibella, 2006), localization of all the three main candidate SOAFs (TR-KIT, PLCZETA, PAWP) seems to be consistent with a possible function in egg activation. However, due to the lack of knock-out mice genetic models, the precise physiological role of TR-KIT, as well as that of other candidate SOAFs, in triggering early embryogenesis remains unknown. It is also possible (and likely) that

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**Table II** Analysis of statistical correlation between seminal parameters and cytofluorimetric data (values = Pearson’s product-moment correlation coefficients)

<table>
<thead>
<tr>
<th></th>
<th>Age</th>
<th>% Clusterin (M3)</th>
<th>% TUNEL</th>
<th>% Double positivity</th>
<th>% TR-KIT</th>
<th>% Atypical forms</th>
<th>Sperm concentration</th>
<th>% Forward motility</th>
<th>Volume of the ejaculate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>0.819</td>
<td>0.964</td>
<td>0.763</td>
<td>0.085</td>
<td>0.921</td>
<td>0.076</td>
<td>0.576</td>
<td>0.076</td>
<td>0.360</td>
</tr>
<tr>
<td>% Clusterin (M3)</td>
<td>-0.19</td>
<td>-0.074</td>
<td>-0.076</td>
<td>-0.076</td>
<td>-0.076</td>
<td>-0.076</td>
<td>-0.076</td>
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<tr>
<td>% TUNEL</td>
<td>0.719</td>
<td>0.799</td>
<td>0.797</td>
<td>0.795</td>
<td>0.791</td>
<td>0.791</td>
<td>0.791</td>
<td>0.791</td>
<td>0.791</td>
</tr>
<tr>
<td>% Double positivity</td>
<td>-0.763</td>
<td>-0.763</td>
<td>-0.763</td>
<td>-0.763</td>
<td>-0.763</td>
<td>-0.763</td>
<td>-0.763</td>
<td>-0.763</td>
<td>-0.763</td>
</tr>
<tr>
<td>% TR-KIT</td>
<td>-0.469</td>
<td>-0.469</td>
<td>-0.469</td>
<td>-0.469</td>
<td>-0.469</td>
<td>-0.469</td>
<td>-0.469</td>
<td>-0.469</td>
<td>-0.469</td>
</tr>
<tr>
<td>% Atypical forms</td>
<td>0.007</td>
<td>0.007</td>
<td>0.007</td>
<td>0.007</td>
<td>0.007</td>
<td>0.007</td>
<td>0.007</td>
<td>0.007</td>
<td>0.007</td>
</tr>
<tr>
<td>Sperm concentration</td>
<td>0.036</td>
<td>0.036</td>
<td>0.036</td>
<td>0.036</td>
<td>0.036</td>
<td>0.036</td>
<td>0.036</td>
<td>0.036</td>
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</tr>
<tr>
<td>% Forward motility</td>
<td>-0.640</td>
<td>-0.640</td>
<td>-0.640</td>
<td>-0.640</td>
<td>-0.640</td>
<td>-0.640</td>
<td>-0.640</td>
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<tr>
<td>Volume of the ejaculate</td>
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<td>-0.235</td>
<td>-0.235</td>
<td>-0.235</td>
<td>-0.235</td>
<td>-0.235</td>
<td>-0.235</td>
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more than a single sperm factor is actually required for full egg activation and progression of early embryonic development.

Regardless of the physiological role played by TR-KIT, however, another important result of our present study is the direct correlation between its presence in human sperm heads and sperm DNA integrity.

TR-KIT expressing spermatozoa were highly enriched in the TUNEL- and clusterin-negative sperm cell populations, whereas spermatozoa with evident morphological anomalies were frequently TR-KIT negative. The percentage of spermatozoa carrying TR-KIT showed a broad variation between different sperm samples from unsolicited volunteer donors. However, almost 100% of the TR-KIT positivity was evident in sperm samples after swim-up selection.

The coincidence of TR-KIT absence and both DNA fragmentation and the high clusterin-positivity, does not appear as a mere consequence of general loss of other proteins in damaged or malformed spermatozoa, since these often retain an intact acrosome in the anterior region of the head, as revealed by the positive staining for acrosin. TR-KIT localization in the perinuclear region of the sperm head might imply an altered formation or damage of the PT as the basis for the correlation between TR-KIT negativity and DNA fragmentation.

Since the TR-KIT expression appears to correlate particularly with sperm DNA integrity, we propose that TR-KIT could be a new marker for evaluating human semen quality, particularly in sperm samples from subjects enrolled in ARTs such as in vitro fertilization (IVF) or ICSI.

Several reports showed no significant correlation between sperm DNA damage and fertilization rates after ICSI (Bungum et al., 2004; Gandini et al., 2004; Greco et al., 2005; Li et al., 2006; Bakos et al., 2008), whereas other reports have shown a negative correlation (Lopes et al., 1998; Benchiba et al., 2003; Huang et al., 2005). These apparent discrepancies could be explained by the technical nature of ICSI, in which morphologically normal sperm cells are selected for injection, thus increasing the chance of using a sperm with intact DNA, by the nature and extent of the DNA damage, and by the variation in DNA repair ability of microinjected oocytes.

Sperm DNA damage might cause arrest of embryonic development at stages in which expression of the paternal genome is required, however impairment of fertilization has been observed even at the level of pronuclei formation (reviewed by Barroso et al., 2009). Since the first steps of embryonic development are mostly under the control of maternal transcripts, these studies suggest that sperm cells carrying DNA damage might also have a lower intrinsic fertilizing capacity.

Our finding that the TR-KIT, a candidate SOAF is absent, or present at lower levels, in DNA damaged spermatozoa, might contribute, at least in part, to their low efficiency in triggering fertilization and/or normal embryonic development after IVF or ICSI reported in some studies.

In conclusion, we have shown that the TR-KIT is present in human ejaculated spermatozoa, where it is localized in the sub-acrosomal and equatorial segment. The equatorial localization of TR-KIT is compatible with its proposed function as a sperm factor at fertilization. Moreover, we found a significant inverse correlation of TR-KIT positivity with markers of sperm DNA damage. Therefore, evaluation of the TR-KIT expression by cytometric analysis might be a predictive parameter of human semen quality, which could be analyzed in hypofertile or infertile subjects. Since we show that high levels of clusterin immunolabeling strongly correlate with DNA fragmentation, our data also suggest that FACS selection with anti-clusterin antibodies of unpermeabilized sperm cells might be used to remove spermatozoa with high levels of DNA damage and low expression of TR-KIT (and, eventually, of other candidate SOAFs which co-localize in the PT).

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**Supplementary data**

Supplementary data are available at http://humrep.oxfordjournals.org/.

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