Identification of pregnancy-associated glycoproteins and alpha-fetoprotein in fallow deer (Dama dama) placenta

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Abstract

Background: This paper describes the isolation and characterization of pregnancy-associated glycoproteins (PAG) from fetal cotyledonal tissue (FCT) and maternal caruncular tissue (MCT) collected from fallow deer (Dama dama) pregnant females. Proteins issued from FCT and MCT were submitted to affinity chromatographies by using Vicia villosa agarose (VVA) or anti-bovine PAG-2 (R#438) coupled to Sepharose 4B gel. Finally, they were characterized by SDS-PAGE and N-terminal microsequencing.

Results: Four distinct fallow deer PAG (fdPAG) sequences were identified and submitted to Swiss-Prot database. Comparison of fdPAG with PAG sequences identified in other ruminant species exhibited 64 to 83% identity. Additionally, alpha-fetoprotein was identified in fetal and maternal tissues.

Conclusion: Our results demonstrate the efficacy of VVA and bovine PAG-2 affinity chromatographies for the isolation of PAG molecules expressed in deer placenta. This is the first report giving four specific amino acid sequences of PAG isolated from feto-maternal junction (FCT and MCT) in the Cervidae family.

Keywords: Affinity chromatography, Fallow deer, N-terminal microsequencing, Pregnancy-associated glycoprotein, Vicia villosa agarose

Background

The family Cervidae includes 40 species and constitutes the second most populous family of artiodactyls. There are four tribes: Cervinae, Muntiacinae, Hydropotinae, and Odocoileinae [1]. Fallow deer belongs to Cervinae tribe, and more specifically to Dama genus. The divergence between Cervidae and Bovidae was estimated to be 22.8 ± 4.7 MYA [2].

In north hemisphere countries, the breeding season of Dama dama occurs throughout October, but may be extended to November. The number of young is one but twins have been observed. The gestation period is 230–240 days; there is no evidence of delayed implantation [3]. The placentation is of oligocotyledonal type, with a maximum of 10 cotyledons in Cervidae family [4]. In histological point of view, placenta from deer is syncytiotrophoblastic [5]. Binucleate cells are a constant characteristic of the trophoblast of the Cervidae. They carry a typical PAS-positive carbohydrate-protein complex [6,7]. They pass from the trophoblast into the crypt lining from the time the villus has occupied a crypt and represent 15–20% of the fetal trophoblastic cells [3].

Pregnancy-associated glycoproteins (PAG) also known as pregnancy-specific protein B (PSPB) or SBU3 antigen constitute a large family of placental glycoproteins [6,8,9]. They are members of the aspartic proteinase gene family and exhibit high sequence identities with other aspartic proteininas, such as pepsinogen, pepsin, chymosin, cathespin D and E [10,11]. Based on their expression throughout the trophoderm and on phylogenetic analyses, the PAG family members are separated into modern (PAG-I) and ancient groups (PAG-II) [12]. The modern PAG are expressed exclusively by binucleate cells whereas the ancient PAG are expressed by both mono- and binucleate trophoblastic cells [13]. The majority of PAG cDNA belongs to PAG-I group [13]. Divergence of PAG-I group is estimated to have taken place 52 ± 6 million years ago. Evolution of the PAG-II group is estimated to have
From the 2000s, it was reported that lectins such as the agglutinins PAG varies from 10 to 17.83% [9, 30]. Indeed, since the early PAG [31]. Therefore, VVA affinity chromatography has been used to wash with 0.9% NaCl and frozen in liquid nitrogen. Placental uterus and placenta tissues were separated, extensively mixed (2 × 3 min) and homogenized (30 min) in 300 ml KCl, pH 7.6) with a ratio 1:3 wt:vol (tissue:buffer). The pH was readjusted regularly to 7.6 with KOH. Phenylmethylsulphonylfluoride (PMSF, 0.2 mM), sodium azide (0.02% NaN₃, wt:vol) and sodium EDTA (0.2% wt:vol) were added at the beginning of mixing. Each homogenate was stirred for 1 h and centrifuged at 20,000 × g during 50 min. The pellets (85.7 and 87.1 g for FCT and MCT, respectively) were taken to a second extraction. They were mixed (2 × 3 min) and homogenized (30 min) in 300 ml and 400 ml of potassium phosphate buffer, respectively. Additional PMSF (0.4 mM), sodium azide (0.04% wt:vol) and sodium EDTA (0.4% wt:vol) were added at the beginning of second extraction. The supernatants issued from the first extraction (0.62 and 1.2 l for FCT and MCT, respectively) were readjusted to pH 7.6 and let stand overnight. The next day, homogenates issued from second extraction were centrifuged (20,000 × g, 50 min). Supernatants from second extraction (0.34 and 0.42 l for FCT and MCT, respectively) were put together with those from first extraction. The pellets (66.7 and 80.4 g for FCT and MCT,

Materials and methods

Collection of cotyledons

Uterus and placenta tissues were harvested from deer (n = 2) during the first half of gestation (110 days post-coitum). Females were taken for slaughter with the agreement of the local ethical authorities from University of Perugia. Immediately after the slaughter of females, uteri and placenta tissues were separated, extensively washed with 0.9% NaCl and frozen in liquid nitrogen.

Measurement of total protein and PAG

Total protein concentrations (TP) of different fractions obtained during the isolation procedure were determined by Lowry method [32], with bovine serum albumine (BSA; ICN Biochemicals Inc., Aurora, OH, USA) as the standard. Due to the absence of specific reagents for PAG measurements in Cervidae species, concentrations of immunoreactive fdPAG were monitored by two different heterologous radioimmunoassay (RIA) systems. Highly purified boPAG67kDa [9] was used as a tracer and standard in both RIA systems. Standard curves ranged from 0.8 to 100 ng/ml. Polyclonal antisera were raised in rabbits (R#) against purified bovine PAG-2 (anti-boPAG-2; R#438) [33–35] and caprine PAG (anti-caPAG55+62kDa; R#706) [22] antigens. Immunisation protocol was previously described by Vaitukaitis et al. [36]. For assay, each fraction was diluted in Tris–HCl buffer (0.025 M Tris, 0.01 M MgCl₂, 0.01% (w/v) sodium azide, pH 7.5) containing 0.1% of BSA. Dilutions ranged from 1:1 until 1:100. Each dilution of the sample (0.1 ml) or standard (0.1 ml) was added to 0.2 ml of assay buffer. Samples were incubated overnight at 20–25°C with 0.1 ml of 125I-PAG (28,000 cpm) and 0.1 ml of each primary antibody (R#438 and R#706 used at initial dilutions of 1:4 000 and 1:120 000, respectively). The total volume of the reaction mixture was 0.5 ml. The next day, 1.0 ml of a double-antibody precipitation system was added to all the tubes except that for total count and a further 30 min incubation took place at room temperature (20–25°C). The end of the procedure was similar to that described previously by Barbato et al. [23].

Isolation of placental proteins

**Protein extraction**

The whole procedure (homogenization, precipitation, centrifugation, and dialysis) was performed at 4°C with the exception of loading and elution of affinity chromatographies (realized at room temperature).

Fetal cotyledonary tissue (FCT; 173.5 g) and maternal caruncula tissue (MCT; 327.5 g) were minced separately. FCT and MCT were mixed five times (5 × 3 min) in potassium phosphate buffer (0.01 M KH₂PO₄ + 0.1 M KCl, pH 7.6) with a ratio 1:3 wt:vol (tissue:buffer). The pH was readjusted regularly to 7.6 with KOH. Phenylmethylsulphonylfluoride (PMSF, 0.2 mM), sodium azide (0.02% NaN₃, wt:vol) and sodium EDTA (0.2% wt:vol) were added at the beginning of mixing. Each homogenate was stirred for 1 h and centrifuged at 20,000 × g during 50 min. The pellets (85.7 and 87.1 g for FCT and MCT, respectively) were taken to a second extraction. They were mixed (2 × 3 min) and homogenized (30 min) in 300 ml and 400 ml of potassium phosphate buffer, respectively. Additional PMSF (0.4 mM), sodium azide (0.04% wt:vol) and sodium EDTA (0.4% wt:vol) were added at the beginning of second extraction. The supernatants issued from the first extraction (0.62 and 1.2 l for FCT and MCT, respectively) were readjusted to pH 7.6 and let stand overnight. The next day, homogenates issued from second extraction were centrifuged (20,000 × g, 50 min). Supernatants from second extraction (0.34 and 0.42 l for FCT and MCT, respectively) were put together with those from first extraction. The pellets (66.7 and 80.4 g for FCT and MCT,
respectively) were submitted to a third extraction in potassium phosphate buffer by using a glass tissue grinder. The two homogenates were centrifuged separately (20 000 × g, 50 min) and the supernatants (390 ml for both FCT and MCT) were added to those from previous extractions. The pellets were discarded.

**Ammonium sulfate (A.S.) precipitation**

The supernatants from the three extractions (FCT or MCT origins) were pooled. They were stirred and dry A.S. was slowly added to obtain 20% saturation solution (113 g/l) (0-20% A.S. fraction). After overnight precipitation, the homogenates were centrifuged at 20 000 × g, during 50 min. The pellets were eliminated. The supernatants (1.32 and 2.0 l for FCT and MCT, respectively) were stirred and dry A.S. was slowly added to obtain 40% saturation solution (121 g/l) (20-40% A.S. fraction). After 3 h precipitation, the homogenates were centrifuged at 20 000 × g, during 50 min and the pellets were discarded. The supernatants (1.34 and 2.1 l for FCT and MCT, respectively) were stirred and dry A.S. was slowly added to obtain 80% saturation solution (281 g/l) (40-80% A.S. fraction). After overnight precipitation, the homogenates were centrifuged at 20 000 × g, during 50 min. The supernatants (1.5 and 2.3 l for FCT and MCT, respectively) were discarded. The pellets (20 g for FCT and 37.2 g for MCT) were diluted in Tris–HCl buffer (0.01 M, pH 7.6) and dialyzed against the same buffer (0.01 M, pH 7.6). The column (8 ml, 2.3 × 2 cm) was equilibrated with the same buffer. After loading, each sample (FCT or MCT) was gently mixed with VVA gel and then incubated overnight at room temperature (RT) into the VVA column. The unbound proteins were washed out with 80 ml of HEPES buffer (0.01 M, pH 7.6). Then, HEPES buffer containing 0.15 M NaCl was loaded onto the column in order to eliminate weaker bound proteins. Proteins were eluted by using the same buffer (0.01 M HEPES + 0.15 M NaCl) added of 0.05 M GalNAc (AppliChem, Darmstadt, Germany). According to their OD, the fractions eluted in the same step (unbound or GalNAc-peak) were pooled, dialyzed (ammonium bicarbonate buffer 0.005 M, pH 8) and lyophilized. VVA gel was regenerated with NaCl (1 M, pH 3) between two consecutive chromatographies.

**Vicia villosa agarose affinity chromatography**

The 40-80% A.S fraction from both FCT and MCT were submitted to R#438 affinity chromatography. Firstly, total immunoglobulin fraction from the immunserum R#438 (lg-438) were purified by ammonium sulphate precipitation and DEAE chromatography [37]. Briefly, 10 ml of crude R#438 were added of 2.5 g of dry A.S. The solution was let stand 20 h at RT. The next day, the solution was centrifuged (10 000 × g, 30 min) and the pellet was washed with 10 ml of 1.75 M A.S. solution. After an additional centrifugation, the pellet was solubilized with 15 ml of distilled water. Precipitated proteins were alternately dialyzed against four batches (5 l) of deionized water and ammonium acetate 0.05 M (pH 5.0). After the last dialysis, proteins were centrifuged (4 000 × g, 20 min) and the supernatant was loaded onto 2.5 ml of DEAE Sephadex A-50 column previously equilibrated with 0.05 M ammonium acetate buffer (pH 5.0). Immunoglobulins were eluted in the non-adsorbed fraction by washing the column with 25 ml of ammonium acetate buffer. Eluted proteins (OD > 0.050) were pooled, dialyzed against 0.005 M ammonium bicarbonate buffer (pH 8) and lyophilized.

Sepharose 4B gel (Amersham Biosciences, Uppsala, Sweden) was activated with cyanogen bromide according to the technique previously described by Axen et al. [38]. Just prior coupled, Ig-438 (30 mg) were solubilized in 15 ml of NaHCO₃ (0.1 M, pH 8.3) containing 0.5 M NaCl. Activated Sepharose 4B gel (10 ml) was added of Ig-438 (30 mg) and stirred 1 h at RT followed by 16 h at 4°C. The next day, unbound Ig-438 was washed out after centrifugation at 1 500 × g (20 min). Unbound sites were blocked by ethanalamine solution (1 M, pH 8). After standing 2 h at RT, the blocking solution was washed away by means of centrifugation. Finally unbound proteins were eliminated by six alternate washes with buffer A (0.1 M sodium acetate adjusted to pH 4 with acetic acid + 0.5 M NaCl) and buffer B (Tris–HCl 0.1 M adjusted to pH 8 ± 0.5 M NaCl).

The Sepharose 4B Ig-438 column (0.7 × 5 cm, 2 ml) was equilibrated with PBS 0.05 M containing 0.15 M NaCl (pH 7.4). A total of 160 and 80 mg of proteins issued from either A.S 40-80% FTC or MCT fractions were loaded three consecutive times. Fractions of 1.5 ml were collected and protein content was monitored by measuring OD at 280 nm. The unbound proteins were eliminated after washing with 20 ml of PBS 0.05 M containing 0.15 M NaCl. In order to ensure the elimination of non-specific weakly bound proteins, a second wash was performed with 20 ml of PBS 0.05 M containing 0.3 M NaCl. Bound proteins were eluted by adding 0.1 M glycine solution adjusted to pH 2.8 with HCl. Before elution, 1.5 ml sodium bicarbonate buffer (0.1 M, pH 8.3) was added to each collection tube.

**Antiserum 438 affinity chromatography**

The 40-80% A.S fraction from both FCT and MCT were submitted to R#438 affinity chromatography. Firstly, total immunoglobulin fraction from the immunserum R#438 (lg-438) were purified by ammonium sulphate precipitation and DEAE chromatography [37]. Briefly, 10 ml of crude R#438 were added of 2.5 g of dry A.S. The solution was let stand 20 h at RT. The next day, the solution was centrifuged (10 000 × g, 30 min) and the pellet was washed with 10 ml of 1.75 M A.S. solution. After an additional centrifugation, the pellet was solubilized with 15 ml of distilled water. Precipitated proteins were alternately dialyzed against four batches (5 l) of deionized water and ammonium acetate 0.05 M (pH 5.0). After the last dialysis, proteins were centrifuged (4 000 × g, 20 min) and the supernatant was loaded onto 2.5 ml of DEAE Sephadex A-50 column previously equilibrated with 0.05 M ammonium acetate buffer (pH 5.0). Immunoglobulins were eluted in the non-adsorbed fraction by washing the column with 25 ml of ammonium acetate buffer. Eluted proteins (OD > 0.050) were pooled, dialyzed against 0.005 M ammonium bicarbonate buffer (pH 8) and lyophilized.

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The Sepharose 4B Ig-438 column (0.7 × 5 cm, 2 ml) was equilibrated with PBS 0.05 M containing 0.15 M NaCl (pH 7.4). A total of 160 and 80 mg of proteins issued from either A.S 40-80% FTC or MCT fractions were loaded three consecutive times. Fractions of 1.5 ml were collected and protein content was monitored by measuring OD at 280 nm. The unbound proteins were eliminated after washing with 20 ml of PBS 0.05 M containing 0.15 M NaCl. In order to ensure the elimination of non-specific weakly bound proteins, a second wash was performed with 20 ml of PBS 0.05 M containing 0.3 M NaCl. Bound proteins were eluted by adding 0.1 M glycine solution adjusted to pH 2.8 with HCl. Before elution, 1.5 ml sodium bicarbonate buffer (0.1 M, pH 8.3) was added to each collection tube.
According to their OD, the fractions belonging the same step (unbound or glycine eluted peak) were pooled together, dialyzed against ammonium bicarbonate buffer (0.005 M, pH 8) and lyophilized.

**Characterization of placental proteins**

**1D-SDS PAGE**

Fractions issued from different fractionation steps were denatured (5 min at 100°C) in Laemmli buffer containing 5% mercaptoethanol. Proteins were separated on a 12% polyacrylamide gel in the presence of SDS on a vertical slab gel system (0.15 × 8 × 7.3 cm). Electrophoresis was performed at 200 V during 40 min. Molecular weight standards (LMW Electrophoresis calibration Kit, Amersham Biosciences, Uppsala, Sweden) were run simultaneously. Proteins were visualized after Coomasie Brilliant Blue R250 staining (Merck, Darmstadt, Germany).

**Western blot**

Details on immunoblotting and Western blot techniques were previously described by Kiewisz et al. [25]. Briefly, proteins were transferred onto a nitrocellulose membrane (0.45 µm, Protran BA85; Schleicher and Schuell Biosciences, Dassel, Germany) after SDS-PAGE. The transfer was performed during 3 h at a constant voltage (60 V) on a TransBlot Cell Apparatus (BioRad, Hercules, CA, USA). Immediately after transfer, membranes were stained with Ponceau Red and extremities of bands corresponding to major proteins were pricked with a needle.

Immunoblotted proteins were probed with two distinct antisera raised against PAG: R#435 (anti-boPAG-2) or R#706 (anti-caPAG55+62kDa). Purification of boPAG-2 and caPAG55+62kDa were described elsewhere [22,33–35]. The final dilution of each first antiserum was 1:100 (0.2 ml antiserum + 4.8 ml PAG-free serum + 15 ml Tris-buffer saline containing 1% BSA).

**Transfer to PVDF membrane and N-terminal microsequence analysis**

Proteins of interest (fractions eluted after VVA or R#438 affinity chromatographies) were separated after SDS-PAGE on a vertical slab gel system (0.1 × 16 × 14 cm; Protein Xi, BioRad). Gels were run at 15 mA/gel during migration in a stacking gel and at 27 mA/gel in the separating gel (12%). Molecular weight standards (LMW Electrophoresis calibration Kit, Amersham Bioscience, Uppsala, Sweden) were run simultaneously. Proteins were transferred onto 0.2 µm polyvinylidene difluoride (PVDF) membranes (BioRad) for N-terminal microsequencing. The transfer was carried out on a TransBlot Cell Apparatus (BioRad) at a constant voltage (60 V) during 2 h 30 min. The PVDF membrane was stained with 0.2% (w/v) Coomassie Brilliant Blue R-250 for 5 min and destained 3 times in 50% methanol solution. After PVDF membranes were dried, proteins were excised and subjected to Edman degradation on a pulsed liquid-phase protein sequencer (Precise 492 Applied Biosystems, Foster City, CA, USA).

The N-terminal sequences obtained in fallow deer have been deposited in the EMBL-EBI database (Swiss-Prot: C0HJC7, C0HJC8, C0HJC9, C0HJD0). N-terminal sequences were compared to those described in databank in order to detect homologies with other isolated native proteins or those deduced from cDNA (Blast, NCBI).

**Results**

**Isolation of fdPAG from FCT and MCT**

After extraction, concentrations of PAG measured by RIA remained very low in both FCT and MCT (data not shown). They were proportionally higher after 3rd extraction than 1st + 2nd extractions. Concerning ammonium sulphate precipitations, as shown in Table 1, in both FCT and MCT tissues, the ratio of equivalent fdPAG measured by using RIA-438 to TP reached higher values after precipitation at 40-80% A.S. saturation. However, when using RIA-706, the ratio of equivalent fdPAG to TP remained very low (<0.06%) in both tissues.

The elution profiles of VVA affinity chromatography are shown in Figure 1A and 1B. After addition of 0.05 M GalNAc, glycoproteins from FCT and MCT were eluted in one major peak.

Finally, with regard to Sepharose 4B Ig-438 affinity chromatography, no protein was eluted after washing with PBS containing 0.3 M NaCl. Addition of glycine-HCl solution (pH 2.8) allowed proteins eluting in one major peak (Figure 2A and 2B).

**Characterization of fdPAG from FCT and MCT**

As shown in Figure 3, after precipitation at 40-80% A.S. of FCT, a single immunoreactive protein (60 kDa) could be observed when using R#435 for Western blot. By using the same antiserum in MCT, two major immunoreactive bands (63 and 66 kDa) were observed after precipitation at 40-80% A.S. saturation. Immunoreactive proteins were also observed in the eluted peak issued from Sepharose 4B Ig-438 affinity chromatography of MCT.

When using R#706, multiple immunoreactive bands (MM ranging from 45 to 70 kDa) could be observed as well after precipitation at 40-80% A.S. saturation than in eluted peaks from VVA and Sepharose 4B Ig-438 affinity chromatographies. Molecular masses were slightly lower after Western blot than after immunoblotting on PVDF membrane and Coomassie staining (58 to 63 kDa) (Figure 4).

Several proteins issued from affinity chromatography and immunoreactive after Western blot with R#435 or R#706 were submitted to N-terminal amino acid sequencing (Figure 4). As described in Table 2, four N-terminal sequences exhibited quite high amino acid identities with...
PAG molecules identified in other ruminant species. These proteins were named according to their molecular masses: fdPAG\(_{62kDa}\)\_A, fdPAG\(_{56kDa}\)\_B, fdPAG\(_{39kDa}\)\_C and fdPAG\(_{56kDa}\)\_D. Moreover, three proteins exhibited high sequence identity with alpha-fetoprotein and another one with albumin.

Sequence comparison of fdPAG (Table 2) with those previously identified in ruminant species ranged from 64 to 83% (Table 3). One PAG sequence exhibited a lower molecular mass (fdPAG\(_{39kDa}\)\_C) and exhibited high amino acid identity with internal fragments of boPAG-6, boPAG-4 and ovPAG-1. The three other sequences appear clearly to correspond to N-terminal PAG termini (fdPAG\(_{62kDa}\)\_A, fdPAG\(_{56kDa}\)\_B and fdPAG\(_{56kDa}\)\_D).

**Discussion**

This paper describes the first isolation and N-terminal microsequencing of PAG molecules from fallow deer. In order to avoid time-related degradation of PAG, our purification protocol was simplified in three main steps: extraction, ammonium sulfate precipitation and affinity chromatographies (VVA or Sepharose 4B Ig-438). As described by Huang *et al.* [27], thawing and freezing moose and elk cotyledons three times and stirring with sand to abrade the surface of binucleate cells was a helpful procedure for PSPB purification. In the present work, the use of tissue grinder during the third extraction of fallow deer tissues was helpful for the recovery of placental proteins from FCT and MCT. Together, these findings

**Table 1 Total protein (TP) and equivalent fallow deer PAG (fdPAG) detected by two heterologous RIA systems**

| Purification step | Fetal cotyledonary tissue | Maternal caruncular tissue |
|------------------|---------------------------|---------------------------|
|                  | fdPAG (mg) and [PAG/TP ratio (%)] | fdPAG (mg) and [PAG/TP ratio (%)] |
| 20-40% A.S.      | 1,401.6 3.16 [0.23]          | 1,956.5 1.23 [0.06]          |
| 40-80% A.S.      | 2,014.9 10.59 [0.53]         | 815.2 3.95 [0.48]           |

*RIA-438, Use of anti-boPAG-2; RIA-706, Anti-caprine PAG\(_{55+62kDa}\).*

**Figure 1** *Vicia villosa* chromatographic profiles from fallow deer. Fractions were issued from fetal cotyledony (A) and maternal caruncula (B) tissues. The column (2.3 x 2 cm; 8 ml) was previously equilibrated with 0.01 M HEPES buffer (pH 7.6). The elution with 0.15 M NaCl or 0.05 M GalNAc buffer (containing 0.15 M NaCl) were designated by arrow. The pooled fractions are in gray.
suggest that in Cervidae species, PAG-PSPB molecules could be strongly related to fetal and maternal membranes. Despite PAG molecules are synthesized in the outer epithelial cell layer (chorion) of fetal cotyledons, purification was performed in both FCT and MCT. Indeed, by using immunocytochemical techniques, PAG-immunoreactivity has been demonstrated not only in fetal cotyledonary but also in maternal caruncular connective tissues in bovine species [14]. Moreover, amount of immunoreactive PAG measured by RIA in *Cervus elaphus* (red deer) extracts was higher in MCT than in FCT [39]. These findings can result from the migration of fetal binucleate cells toward the maternal junction.

Low PAG concentrations were observed in both FCT and MCT extracts (use of heterologous RIA-438 and RIA-706). In the same way, concentrations of PAG measured in plasma from pregnant fallow deer are very low [40–42]. Low concentrations can be related to the use of a bovine PAG-PSPB preparation as standard and tracer. According to Brandt et al. [28], PAG from white-tailed deer are mostly from PAG-II group. This observation was confirmed by results presented in Table 1, which reports that immunoreactivity of fdPAG is much higher when measured by using the anti-boPAG-2 antiserum (R#438).

Molecular masses of three fdPAG (calculated after Coomassie staining) ranged from 56 to 62 kDa. They are in the range of those described in other species [43] and are slightly lower than those observed after Western blot. In previous works, the same phenomenon was observed in both buffalo [23,24] and American bison [25]. Unfortunately, the explanation of such a difference is not known. However, it can be hypothesized that the calculation of molecular mass after Western blot is less precise due to the greater width of bands. Accordingly, we choose to assign proteins on the basis of their molecular mass calculated after Coomassie blue staining.

Interestingly, N-terminal extremities of fdPAG_{62kDa_A} and fdPAG_{56kDa_B} did not show the highly conserved RGS- amino acid residues previously reported for other ruminant species [19–26]. However, they fit with the

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**Figure 2** Ig-438 affinity chromatography profiles from fallow deer. Fractions were issued from fetal cotyledonary (A) and maternal caruncula (B) tissues. Column (0.7 × 5 cm, 2 ml) was previously equilibrated with PBS 0.05 M containing 0.15 M NaCl (pH 7.4). Unbound proteins were eliminated after washing with PBS 0.05 M containing 0.15 M NaCl, followed by washing with PBS 0.05 M containing 0.3 M NaCl. Bound proteins were eluted by adding 0.1 M glycine solution adjusted to pH 2.8 with HCl. Before elution, 1.5 ml sodium bicarbonate buffer (0.1 M, pH 8.3) was added to each collection tube.
PGS- sequence described for Deer PAG-1 and −2 in white-tailed deer [28]. In their sequenced part, identities of proteins isolated in fallow deer and identified in white-tailed deer ranged from 67 to 80%. This discrepancy can be partially explained by the phylogenetic divergence between *Dama dama* and *Odocoileus virginianus*, estimated to be 7.17 MYA [44].

Concerning belonging of fallow deer PAG to modern (PAG-I) or ancient (PAG-II) groups, as described in Table 3, N-termini of fdPAG62kDa_A and fdPAG56kDa_B exhibit homologies higher than 70% with different proteins from PAG-I group previously identified in white-tailed deer [28], cow [9,10], and ewe [10] placentas. On the other side, fdPAG56kDa_D N-terminal sequence was identified as relatively close to boPAG-12 (PAG-II group). Nevertheless, as N-terminal micro-sequencing only refers to a limited number of residues, we could not perform the comparison of the whole sequences from *Dama dama* with PAG molecules deduced from cDNA and described in other ruminant and porcine species [43].

An additional protein (fdPAG39kDa_C) issued from VVA chromatography (FCT) exhibited a lower molecular mass. This protein showed a high amino acid identity with internal fragments of boPAG-6, boPAG-4 and ovPAG-1, indicating that it corresponds to an internal fragment of PAG. A similar finding was described by Doré *et al.* [45] concerning the porcine basic protein, which corresponds to an internal fragment of porcine PAG.

Huang *et al.* [27] described the use of an affinity chromatography developed by antiserum anti-bovine PSPB for isolation of PSPB molecules from moose and elk placenta. However, they did not characterize the proteins they could obtain. In the present study, we performed affinity chromatographies on FCT and MCT by using an antiserum

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**Figure 3** Western blot of fdPAG proteins extracted from fetal cotyledonal (FCT) and maternal caruncular (MCT) tissues. Two polyclonal antiseras raised against boPAG-2 (AS#435) or caprine PAG55+62kDa (AS#706) were used as primary antiserum at 1:100 dilution. Molecular weight standards (kDa, 7 μg/lane) were loaded on the right or left position of figures. Lane 1: A.S. 40-80% fraction from FCT; Lane 2: VVA eluted peak from FCT; Lane 3: eluted peak from Sepharose 4B Ig-438 affinity chromatography from FCT; Lane 4: A.S. 40-80% from MCT; Lane 5: VVA peak from MCT; Lane 6: eluted peak from Sepharose 4B Ig-438 affinity chromatography from MCT. Fifty μg were loaded in lanes 1 and 4; 30 μg were loaded on lanes 2, 3, 6 and 7.
raised against a boPAG-2 preparation [33-35]. Western blot analysis highlighted immunoreactivity of several proteins issued from Sepharose 4B Ig-438 affinity chromatography with both R#435 and R#706. This chromatography allowed the successful identification of one fdPAG while three other sequenced proteins corresponded to alpha-fetoprotein (AFP, also known as fetuin A or alpha-2HS glycoprotein). AFP is the major serum protein in fetal ruminants, pigs, as well in humans and rodents [46]. This protein was firstly described in 1944 by Pedersen [47] and further characterized by Spiro [48]. AFP is synthesized and secreted by the fetal liver [49] and to a lesser extent the placenta, kidneys and the tongue [50]. Molecular mass of AFP is very similar to those from PAG-PSPB molecules. It ranges from 51 to 67 kDa depending on carbohydrate content (6 to 8%). However, in contrast to PAG-PSPB, the concentrations of AFP do not increase in maternal circulation during gestation [49] and cannot be used for pregnancy diagnosis in cattle [51,52].

It is noteworthy that Butler et al. [8] co-purified AFP and PAG-PSPB from bovine placental tissues collected between Days 16 and 280. Thus, it cannot be excluded that boPAG-2 antigen (used to generate R#438) could be contaminated by AFP. We hypothesize that contamination of boPAG-2 preparation with AFP may explain at least partially the poor ability of Sepharose 4B Ig-438 affinity chromatography to isolate PAG in fallow deer species. However, other factors such as phylogenetic divergence between PAG molecules isolated in fallow deer and bovine species cannot be excluded.

### Table 2 Molecular masses and N-terminal sequences of proteins isolated from fallow deer (Dama dama) placenta

| Origin of tissue | Affinity chromatography used for isolation | MM (kDa) | Sequenced fragment* | Protein | Accession number |
|------------------|------------------------------------------|---------|---------------------|---------|-----------------|
| FCT, Fetal cotyledonary tissue | WA | 62 | YQKSSPGSNITT | fdPAG_{\text{62kDa}}A | C0HJC7 |
| | | 56 | YQTSSPGSNITGL | fdPAG_{\text{56kDa}}B | C0HJC8 |
| | | 39 | DVGPSTLANN | fdPAG_{\text{39kDa}}C | C0HJC9 |
| Antiserum R#438 Sepharose | | 56 | IPLDITAGYKE | Alpha-fetoprotein | C0HJD1 |
| MCT, Maternal caruncular tissue | WA | 62 | DTHKSEIAHR | Albumin | C0HJD2 |
| Antiserum R#438 Sepharose | | 63 | IPLDITAGY | Alpha-fetoprotein | C0HJD1 |
| | | 60 | IPLDITAGYKE | Alpha-fetoprotein | C0HJD1 |
| | | 56 | SLRKMHALGET | fdPAG_{\text{56kDa}}D | C0HJD0 |

FCT, Fetal cotyledonary tissue; MCT, Maternal caruncular tissue; MM, Molecular mass.

* Amino acid microsequence analysis was performed by the automated Edman degradation method.

**Figure 4** Coomassie blue stained PVDF membrane after SDS-PAGE. Low molecular weight standards (kDa; 7 μg/lane) were loaded on the right and the left side of the figure. Lane 1: eluted peak from Sepharose 4B Ig-438 affinity chromatography from FCT; Lane 2: VVA eluted peak from FCT; Lane 3: eluted peak from Sepharose 4B Ig-438 affinity chromatography from MCT; Lane 4: VVA peak from MCT. Fifty to 75 μg were loaded in each lane.
Regarding VVA affinity chromatography, the major protein obtained from MCT corresponded to the N-terminal sequence of serum albumin. Both albumin and AFP belong to albuminoidal gene superfamily. They are known to bind and/or transport a multitude of ligands, such as bilirubin, fatty acids, steroids, heavy metals and others [53]. Mature bovine serum albumin (BSA) is a single chain non-glycosilated polypeptide (583 amino acids long; accession number P02769) and contains three structural domains. BSA displays a molecular mass of 66 kDa. In fallow deer, apparent molecular mass was slightly lower after SDS-PAGE (62 kDa). As serum albumin is the most abundant protein of the circulation [54], it is hypothesized that fallow deer albumin detected in extracts of extraembryonic membranes might have been due to its higher abundance in maternal blood within the cotyledons.

When data of Table 3 are considered, it appears that albumin shows the same N-terminal sequence in cow, goat and fallow deer, whereas alpha-fetoprotein and PAG exhibits lower identities with other ruminants (82 to 91% and 64 to 83%, respectively). This observation can be related to the rapid evolution of PAG having lead to the high diversity observed in ruminant species [11,12,55].

Concerning biological aspects of PAG, an interesting review by Roberts et al. [55] suggested that PAG could sequester or transport small peptides in their binding cleft (6–8 amino acids long) [11]. An alternative hypothesis was recently described by Telugu et al. [56] who reported that placental aspartic proteinases might participate in placental remodeling by means of proteolytic digestion of endocytosed proteins in the uterine lumen. In the present study, N-terminal sequences of PAG extracted from Dama dama exhibited high identities with those from both modern (PAG-I, incapable of acting as proteolytic enzymes) and ancient groups (PAG-II, predicted to possess proteolytic activity). Moreover, our investigation confirmed that deer PAG molecules require abrasive extraction from cotyledonary tissue. Together, these findings claim for PAG as participating to a complex network of tight-junction-associated proteins communicating at feto-maternal interface.
Conclusion
This is the first study describing the isolation and characterization of PAG from fallow deer placenta. The use of VVA and Sepharose 4B Ig-438 affinity chromatographies allowed the identification of four PAG molecules. Analysis of N-terminal sequences revealed high sequence identity with PAG from other ruminant species.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
MB performed experimental work, data analysis and drafted the manuscript. AFT assisted in the design of study and participated in carrying out chromatography and radioimmunoassay. OB participated in the design of the study. JFB conceived the design of the study, coordinated the work and helped in writing the manuscript. NMS participated in carrying out PAG purification and sequence analysis, and had important input into and participation in writing the manuscript. All authors read and approved the final version of the manuscript.

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