The evolution of interferon-tau

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Abstract

Thirty years ago, a novel type I interferon (IFN) was identified by molecular cloning of cDNA libraries constructed from RNA extracted from ovine and bovine pre-implantation embryos. This protein was eventually designated as IFN-tau (IFNT) to highlight its trophoblast-dependent expression. IFNT function is not immune related. Instead, it interacts with the maternal system to initiate the establishment and maintenance of pregnancy. This activity is indispensable for the continuation of pregnancy. Our review will describe how IFNT evolved from other type I IFNs to function in this new capacity. IFNT genes have only been identified in pecoran ruminants within the Artiodactyla order (e.g. cattle, sheep, goats, deer, antelope, giraffe). The ancestral IFNT gene emerged approximately 36 million years ago most likely from rearrangement and/or insertion events that combined an ancestral IFN-omega (IFNW) gene with a trophoblast-specifying promoter/enhancer. Since then, IFNT genes have duplicated, likely through conversion events, and mutations have allowed them to adapt to their new function in concert with the emergence of different species. Multiple IFNT polymorphisms have been identified in cattle, sheep and goats. These genes and gene alleles encode proteins that do not display identical antiviral, antiproliferative and antiluteolytic activities. The need for multiple IFNT genes, numerous alleles and distinct activities remains debatable, but the consensus is that this complexity in IFNT expression and biological activity must be needed to provide the best opportunity for pregnancy to be recognized by the maternal system so that gestation may continue.

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Introduction

In nearly every eutherian mammal, pregnancy can only be maintained if progesterone production continues beyond the length of a normal estrous or menstrual cycle. This requires that the ovarian structure producing progesterone, termed the corpus luteum (CL), must remain active either throughout gestation or until the placenta produces sufficient progesterone. Rescuing the CL is achieved in various ways. Higher primates, for example, produce chorionic gonadotrophin (CG; hCG in humans) to maintain CL function. Sheep, cattle, goats and presumably other pecoran ruminants rescue CL function by preventing the pulsatile release of uterine prostaglandin F2α (PGF2α), the luteolytic agent that causes CL regression (Spencer et al. 2004, Roberts et al. 2008). This event is one critical feature of physiological adjustments the maternal system must make in order to accommodate pregnancy. These maternal adjustments to pregnancy are generally referred to as maternal recognition of pregnancy.

Moor and Rowson initially investigated maternal recognition of pregnancy in ruminants by completing embryo recovery and transfer studies in sheep and cattle (Moor & Rowson 1964, 1966, Rowson et al. 1969, Lawson et al. 1975). This work defined the critical window of pregnancy recognition in these species and suggested the active factor was proteinaceous. The protein inferred responsible for this activity was given a number of names, which included trophoblastin, protein-X and trophoblast protein-1 (oTP1 and bTP1 for ovine and bovine counterparts respectively) (Martal et al. 1979, Godkin et al. 1982, 1984a). Functional studies also were completed, where either protein extracts from pregnant uterine flushes or purified proteins from short-term ovine conceptus cultures extended CL function for several days to a few weeks (Rowson & Moor 1967, Martal et al. 1979, Godkin et al. 1984b). This antiluteolytic activity and various additional pregnancy-based biological activities for IFNT have been described in several recent reviews, including three in this special issue (Spencer & Hansen; Forde & Lonergan; Bazer & Thatcher).

In 1987, cDNA screening and sequencing using an ovine conceptus cDNA expression library and oTP1 antibody uncovered this maternal recognition of pregnancy factor as a type I IFN (Imakawa et al. 1987). Soon thereafter, cDNA screening for the sequence encoding ovine trophoblastin revealed that it represented the same protein (Charpigny et al. 1988). Screening a bovine cDNA conceptus library determined that bTP1 also was a type I IFN (Imakawa et al. 1989). At the time when the identity of IFNT was revealed, IFNs were thought of solely as antiviral and immune-regulatory agents. The IFN field had developed to the point where...
numerous IFNs had been well established as antiviral agents and were being considered as therapies for various viral infections and cancers (Pestka et al. 1987). It was a considerable surprise when the pregnancy recognition factor for ruminants was identified as a type I IFN. In fact, this finding was challenged, arguing that a type I IFN could not function as an essential hormone of pregnancy because fertility was not compromised in mice lacking components of IFN signaling (Roberts et al. 1998). Much has been learned since this point, including that the mouse cannot be used as model to study pregnancy recognition in ruminants.

This review will describe how this unique subclass of IFNs emerged from other type I IFNs. We also will speculate why IFNT was selected to serve as the maternal recognition of pregnancy factor in ruminants. Several excellent reviews already exist on this topic (Roberts et al. 2003, 2008). This article will summarize that information and provide additional insights into what has been learned and what still remains unclear about the evolution of IFNT.

Interferons

Three main subclasses of IFNs exist. These are classified as type I, II or III IFNs. The type I IFNs are composed of at least 9 distinct subfamilies: IFN-alpha (IFNA), beta (IFNB), delta (IFND), omega (IFNW), epsilon (IFNE), kappa (IFNK), tau (IFNT), zeta (IFNZ) and X (IFNX) (Krause & Pestka 2005, Walker & Roberts 2009). The type I family likely emerged from the type III IFN family, which is described below, as mammalian lineages diverged from avians and reptiles ~300 million years ago (Hughes 1995, Levraud et al. 2007). Seven of these nine IFN subfamilies have been identified in the bovine genome (IFNA, IFNB, IFNE, IFNK, IFNW, IFNT, IFNX) (Walker & Roberts 2009). The selectivity of most IFN responses are governed by tissue-dependent expression and type of pathogen encountered. The IFNA and IFNB subfamilies are often referred to as immediate viral responders. Many cell types produce IFNA in response to various stimuli, whereas IFNB is primarily secreted from fibroblasts (Pestka et al. 2004). Most of the remaining type I IFNs appear to have more specialized niches, either by their production in specific cell types or by their involvement with reacting to specific types of viruses (Pestka et al. 2004).

Two IFNs, IFND and IFNT, are not regulated principally by responding to viral infections but rather are expressed constitutively by pre-implantation conceptuses of pigs and ruminants respectively. IFND is not responsible for pregnancy recognition in pigs, but it may have important auxiliary roles in maintaining pregnancies (Lefevre et al. 1998a,b). IFND emerged in mammal genomes much earlier than IFNT, and it is not a direct ancestor of IFNT (Cochet et al. 2009). Noteworthy features about IFND gene retention in mammals and its potential biological functions during pregnancy in pigs have been described elsewhere (Roberts et al. 2008, Sang et al. 2014).

All type I IFNs exert their actions through interaction with the same receptor complex (Roberts 2007, Roberts et al. 1992a). These receptors are termed IFNA receptors (IFNAR). They are composed of two heterologous subunits, termed IFNAR1 and IFNAR2. Receptors mediate transcription by controlling the activity of Janus kinases and signal transducers of transcription (JAK-STAT) (Roberts 2007). These receptors confer all the activities for type I IFNs throughout the body, including the IFNT signals generated within the epithelial and stromal endometrium in ruminants (Li & Pestka 1994a).

Type II IFN is represented by a single IFN, known as IFN gamma (IFNG). IFNG contains antiviral, antitumor and immune-modulating activities and is required for innate and adaptive immunity against viral and bacterial infections, but it is best known as an inflammatory and autoimmune cytokine (Bach et al. 1997, Billiau & Matthys 2009). IFNG has no structural similarity with type I or III IFNs. It is derived from a single gene, functions as a homodimer and acts through a receptor complex distinct from those utilized by type I IFNs (Billiau & Matthys 2009).

Less is known about the type III IFN family due, in part, because members of this grouping are not present in all mammals (Galani et al. 2015). There are three members of this family in humans. These are known either as IFN-lambda (IFNL) 1, 2 and 3 or interleukin (IL) 29, 28A and 28B. These IFNs share low structural similarity with type I IFNs and IL10 (Gad et al. 2009). They utilize a receptor complex that differs from that used by type I IFNs (Sheppard et al. 2003). This receptor complex contains a specific, unique alpha subunit and a common IL10 beta subunit. This class of IFNs likely only plays a minor role in controlling viral infections (Ank et al. 2008), although it has been implicated in the viral defenses of the human placenta (Bayer et al. 2016).

The origins of IFNT

All type I IFN genes reside within a specific locus on chromosome 8 in cattle (Walker & Roberts 2009). IFNT shares its greatest sequence identity with IFNW (~70% amino acid (AA) identity) and contains only moderate similarity with IFNA (~50% AA identity) and low similarity with IFNB (~25% AA identity) (Leaman et al. 1992, Roberts et al. 1992b). Although the primary AA sequences of type I IFNs differ, each of these IFNs possesses a similar secondary and tertiary structure. Conserved cysteine residues exist in all IFN subclases, and disulfide bonding and conservation of five alpha helices arranged in an antiparallel manner and connected by definitive loop regions produces similar
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Genus/species: *Bos taurus* (Cow), *Bubalus bubalis* (Water buffalo), *Capra hircus* (Goat), *Cervus elephus* (Red deer), *Giraffa camelopardalis* (Giraffe), *Moschus berezovskii* (Musk deer, dwarf), and *Ovis aries* (Sheep).

The entire coding region sequences (signal peptide plus mature protein) were analyzed using Molecular Evolutionary Genetics Analysis version 7 software (MEGA7) (http://www.megasoftware.net/). Maximum Likelihood analysis was completed using the Bootstrap method with 2000 replicates using the Tamura-Nei model (Tamura & Nei 1993). Trees were developed using the Interactive Tree of Life version 3 software (Letunic & Bork 2016).

The functional significance of the 5' UTR of this gene may restrict the functional expression of the gene. A phylogenetic tree (Fig. 1) shows the probable evolutionary relationships between representative *IFN* genes from several ruminant species and representative *IFNTA*, *IFNB*, and *IFNW* from the cow, sheep, and goat. Clustering of the four major *IFN* classes is evident, with one exception. Gayal *IFNT* (*Bos frontalis*, Mithun) associates more closely with *IFNW* genes instead of the remaining *IFNT* genes. There is low confidence in where this specific *IFNT* should be placed. Bootstrap values, which indicate the probability of proper placement within the phylogenetic tree, offer a low statistical likelihood of its exact location. Usually bootstrap values >0.7 is needed to have confidence that genes are inserted in the proper location within the tree.

A phylogenetic tree (Fig. 1) shows the probable evolutionary relationships between representative *IFNT*-coding region nucleotide sequences from several ruminant species and representative *IFNTA*, *IFNB*, and *IFNW* from the cow, sheep, and goat. Clustering of the four major *IFN* classes is evident, with one exception. Gayal *IFNT* (*Bos frontalis*, Mithun) associates more closely with *IFNW* genes instead of the remaining *IFNT* genes. There is low confidence in where this specific *IFNT* should be placed. Bootstrap values, which indicate the probability of proper placement within the phylogenetic tree, offer a low statistical likelihood of its exact location. Usually bootstrap values >0.7 is needed to have confidence that genes are inserted in the proper location within the tree.

Usually bootstrap values >0.7 is needed to have confidence that genes are inserted in the proper location within the tree. The bootstrap value for Gayal *IFNT* is 0.3 (Baldauf 2003). Also, this sequence was derived from genomic DNA, and its tropoblast-dependent expression remains uncertain (Rajaravindra et al. 2006). Therefore, this gene may well be an *IFNW*, of which there are >20 members in the American bison gene may reflect inter-genus divergences. The present-day bison genome contains bovine ancestry because of accidental and intentional matings over the past several hundred years (e.g. Beefalo) (Hedrick 2009). Only 1.5% of all plains bison are predicted to be free of domestic cattle ancestry (Freese et al. 2007). The inter-genus association between the water buffalo *IFNT* sequence with the goat sequence cannot be explained. The presence of *IFNT* genes in the final ruminant family, the Antilocapridae, has not been investigated. The pronghorn is the only living species in this family.

The main distinguishing feature between *IFNT* and most other type I *IFNs* is its unique, tropoblast-specific expression in ruminant species. The first 400 bases upstream of the *IFNT* gene transcription start site are unique to *IFNT*, and this region lacks any apparent virus-inducible transcriptional elements (Leaman et al. 1992). Several tropoblast-specifying transcriptional control elements exist within this 5' promoter/enhancer region, including ETS2 and DLX3 (Ezashi et al. 1998, Imakawa et al. 2006). This unique 5' UTR may restrict *IFNT* transcription to tropoblast cells, the outermost cells of a developing conceptus that will develop into the outermost layer of the placenta. In bovine embryos, expression begins as the tropoblast lineage emerges at the morula and blastocyst stages (day 6–7 of pregnancy in cattle) and continues until tropoblast cells begin adhering to the uterine epithelium (~day 16 in cattle) (Ealy et al. 2001, Kubisch et al. 2004). The production of *IFNT* is small at the morula and blastocyst stages but increases as conceptus elongation occurs (day 14–17 in cattle). The control of *IFNT* expression is covered in greater detail in a companion review article (Ezashi & Imakawa).

Another distinguishing feature of *IFNT* is its unique 3' UTR. The proximal 300 bases of the 3' UTR are highly conserved in *IFNT* (Leaman et al. 1992). The conservation with *IFNW* sequences ends approximately 120 bases downstream of the translational stop codon (Leaman et al. 1992). The functional significance of this unique 3' UTR is unclear. At least two micro-RNA (miRNA) target sequences exist within the 3' UTR in bovine and ovine *IFNT*, and at least one of these miRNA is differentially expressed in bovine conceptuses during early pregnancy (Yang & Ealy Unpublished observations).
However, it is not clear if miRNAs exert any control on IFNT production in ruminants.

Based on the observations that IFNT is similar in coding sequence to IFNW but different in its 5’ and 3’ UTRs, we predict that the ancestral IFNT gene was created from rearrangement or insertion events, where an IFNW gene was combined with a novel, trophoblast-specifying 5’ UTR and a distinct 3’ UTR. The timing of these rearrangement events is predicted to have occurred around 36 million years ago (MYA) (Roberts et al. 1997), which coincides with the deviation of Ruminantia suborder from Suiforme (swine) and Tylopoda (llamas, camels) suborders of Artiodactyla (Roberts et al. 1998). Initial indications that IFNT was restricted to ruminant species were observed by using Southern blot analysis (Fig. 2) (Leaman & Roberts 1992). Use of a 3’ UTR ovine IFNT probe to distinguish IFNT from what we now know to be numerous IFNW provided the first evidence that IFNT was unique to the ruminant species (Fig. 2). In most cases, multiple bands were detected within these species, while there was no hybridization to DNA from other mammalian taxa. Subsequent whole genome sequencing has confirmed this ruminant-specific inference.

![Figure 2](https://www.reproduction-online.org/)

**Figure 2** Genomic Southern Blot analysis of gene distribution for IFNT genes in various species (Zoo Blots). DNA (3–8 μg) from indicated species was digested with EcoR1, electrophoresed, transferred to a nylon membrane, and hybridized with an O. aries IFNT probe derived from the 3’ UTR region. Molecular sizes are indicated in kilobase (kb) pairs. This figure was taken from (Leaman et al. 1992).

**Interferons at the placental–maternal interface**

The evolution of maternal recognition of pregnancy signals has not been defined in many species, but what is known is that these signals usually are derived from the trophoblast cell, are highly variable, and they appear to be evolving rapidly (Roberts et al. 2016). For example, higher primates and Suinae, both of which lack IFNT, utilize CG and estrogens respectively, as their pregnancy recognition factors. The early timing of implantation and extensive eroding of the endometrium for intimate access to maternal blood permits hCG to act as an endocrine-mediated luteotrophic factor for humans. By contrast, no endometrial erosion occurs in the pig placenta, and estrogen must diffuse into the maternal bloodstream. Estrogens also function locally within the uterus to prolong CL lifespan (Bazer 1989, Ziecik 2002).

Maternal recognition of pregnancy is very different in ruminants. One notable feature of conceptus development in these species is that implantation is delayed until after maternal recognition of pregnancy has taken place. Like other ungulates (Perisodactyla and Artiodactyla), ruminant conceptuses remain free-floating after hatching from the zona pellucida (day 8–10 of pregnancy in cattle). Intimate contact with the uterus does not occur until on or after day 16 of pregnancy in cattle (Bazer et al. 2009). This requires the development of a very different pregnancy signal, one that can function locally within the uterus instead of systemically. The evolution of IFNT has satisfied this role in ruminants, but other ungulates utilize different signals. As mentioned above, the pig utilizes estrogens secreted from trophoblast cells during early pregnancy as their antiluteolytic signal (Bazer 1989). The pregnancy recognition signals utilized by most other ungulates have not been described, but those that have been studied thus far, such as the horse (Klein & Troedsson 2011), not utilize either an IFN or a steroid hormone to confer pregnancy recognition.

The selective pressure to utilize such unexpected and unique factors as maternal recognition of pregnancy signals likely occurs because of selective pressures placed onto mammals to diversify placental types. Maternal control of metabolic resource management is one of the best explanations for why so many different placental types exist in mammals (Fowden & Constancia 2012). This is one aspect of what is conventionally termed the genetic conflict theory, where females seek to balance available resources with reproduction to maximize the dissemination of their genetics to future generations (Crespi & Semeniuk 2004). The vast array of placental types that exist in mammals likely represents the various ways that maternal and fetal systems have evolved to provide sufficient maternal resources without compromising maternal health and survival. Another probable reason for placental evolution is for avoidance of immune rejection (Hemberger 2013). Processes
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The multiplicity of IFNT genes and alleles

Cattle contain at least 13 IFNA, 6 IFNB and 24 IFNW genes (Walker & Roberts 2009). The exact number of IFNT genes in ruminants remains debatable, but multiple genes are certain when examining genomic DNA hybridization studies (Fig. 2) and more recent genome sequencing analyses (Walker & Roberts 2009). Only three IFNT genes are annotated in the current bovine genome assembly (UMD 3.1.1). We contend that additional genes will be detected as the bovine genome becomes more completely annotated. A careful examination of the bovine genome revealed 45 distinct IFNT sequences (Walker & Roberts Unpublished observations). The ~7-fold depth of coverage for this genome would place the average genome read number at between 6 and 7 IFNT genes. Supportive evidence of this is seen when examining transcript variants from individual conceptuses (Walker & Roberts 2009). At least 6 different IFNT variants are expressed in individual bovine conceptuses and from a bovine trophoblast cell line (CT1) derived from a single blastocyst. Some of these variants may be allelic. It is also possible that the number of IFNT genes may vary among individuals and breeds.

There are many examples of gene family expansions, but the process appears to be particularly common for families whose members are expressed in the placenta (Rawn & Cross 2008, Roberts et al. 2016). Examples include (1) growth hormone-related and prolactin-related proteins found in primates, rodents and ruminants, (2) the CG beta gene family in primates, (3) pregnancy-specific proteins found in primates and rodents, (4) pregnancy-associated glycoproteins found in ruminants and swine and (5) IFNT in ruminants. Having several IFNT genes may provide extra expression when a massive boost in IFNT is needed when the CL of pregnancy is hovering on the brink of luteolysis. On the other hand, duplication will ultimately lead to redundancy, such that unnecessary genes can be simply eliminated without consequence. Occasionally, however, such redundancy may allow a duplicated gene time to acquire mutations that improve old functions and even acquire new ones (Hughes 1995). A similar process operating on an ancestral IFNW probably allowed such a gene to acquire trophoblast-specific regulatory regions and evolve into what we now know as IFNT.

Several IFNT gene variants have been described in cattle, sheep and goats. It is not clear whether these variants represent individual genes or allelic variants of a few genes. Figure 3 illustrates a phylogenetic tree that includes every unique IFNT sequence deposited in the GenBank database for the cow, sheep and goat. This analysis predicts how variants likely evolved from each other based on the number and location of individual nucleotide changes. Nucleotide sequences

Figure 3 Phylogenetic analysis of nucleotide coding sequences for all polymorphic forms of IFNT available in the GenBank database for cattle (B. taurus), sheep (O. aries) and goat (C. hircus). The identity of each gene used for this analysis is provided in Supplementary Table 2. The coding regions without their signal peptide regions (i.e. mature peptide only) were analyzed using Molecular Evolutionary Genetics Analysis version 7 software (MEGA7) (http://www.megasoftware.net/). Maximum Likelihood analysis was completed using the Bootstrap method with 2000 replicates using the Tamura-Nei model (Tamura & Nei 1993). Trees were developed using the Interactive Tree of Life version 3 software (Letunic & Bork 2016). The Bos taurus IFNT1, IFNT2 and IFNT3 genes are represented by cow sequence# 9, 1 and 6 respectively. The Ovis aries IFNT1 and IFNT2 genes are represented by sheep sequence# 5 and 9 respectively. The Capra hircus IFNT gene is represented as goat sequence# 1.
were examined rather than AA sequence divergence in order to obtain a more complete picture of IFNT gene variant evolution and to strengthen their correct placement within the tree (Baldauf 2003). Since an exact identity of these variants is not required for this discussion, variants were given arbitrary numbers to identify them (Supplementary Table 2 for the GenBank accession number key, see section on supplementary data given at the end of this article). Eighty-one sequences were evaluated; 29 in the cow, 23 in the sheep and 29 in the goat. Eighteen bovine, 12 ovine and 10 caprine IFNT variants were validated by their detection in multiple RT-PCR reactions from identical samples. These observations rule out the possibility that the nucleotide changes were acquired during amplification. The remaining variants have not gone through this validation process, and so, it is not certain if their mutations are represented in the cow, sheep and goat genomes or if they represent mutations introduced during RT or PCR. Their inclusion was chosen in order to present the most extensive depiction of IFNT polymorphisms.

Clear distinctions between the three species are evident. Only two variants did not segregate within their respective species (sheep 14 and goat 23). The most obvious example of gene clustering occurred in the goat, where major groups of transcripts exist. Several smaller clusters of IFNT variants were also seen within these larger goat clusters. Clustering also occurred in the sheep and cow variants. Some clusters of IFNT variants contain only a few members, whereas other clusters contain many. No breed-dependency is apparent for these alleles in any species, but further work is needed to explore this possible explanation for the multitude of polymorphisms. It also is interesting to note that most IFNT variants have appeared recently in ruminants (Fig. 3). The rapid timescale for the evolution of these expansive clusters of related IFNT variants are suggestive of gene conversion events, where crossover recombination events transfer genetic material from one chromosome to the other (Cole et al. 2012). In the case of intron-less genes like IFNT and other type I IFNs, the entire gene can be recombined to generate highly similar but novel alleles.

Only a subset of these nucleotide polymorphisms will modify the primary AA sequence of IFNT. Bovine IFNT variants have been the best studied in this regard (Walker & Roberts 2009). In most gene variant to variant comparisons, the incidence of synonymous nucleotide changes (no change in AA codon read) usually is either equal to or greater than the rate of acquisitions of non-synonymous changes (change in codon read). This suggests that the open reading frames of the IFNTs are no longer under strong selective pressure to change primary AA sequence to greatly affect potency or stability. Rather, most of the variation between IFNT isoforms appears to be consistent with purifying selection.

**Why are there so many IFNT genes and gene variants?**

The reason why there is a need for several IFNT genes in ruminants has been the topic of much research over the past 20 years. Several theories have been put forth. As noted in the previous section, multiple genes may exist to maximize IFNT production during early pregnancy. IFNT is the most abundant transcript and protein during the peri-implantation period. Hundreds of micrograms of IFNT is produced by a single elongated conceptus during peak production (Godkin et al. 1982, Ashworth & Bazer 1989). Alternatively, perhaps multiple genes are needed to ensure adequate coverage of IFNT activity as conceptuses develop and circumstances change in the uterus as it prepares for placental attachment. In cattle, two IFNT genes are transcribed more highly than other genes at the blastocyst stage, but the expression of several additional IFNT variants are upregulated as conceptus development advances (Walker & Roberts 2009). Also, differences exist in the abundance of at least two ovine IFNT transcripts when conceptuses are undergoing implantation (Sakurai et al. 2013).

Another reason why multiple IFNT genes may exist is that they encode for proteins with different biological potencies. The polygenic nature of type I IFNs as antiviral agents has afforded IFNs with the ability to evolve and adapt to their ever-changing environment and viral insults (Goodbourn et al. 2000). Perhaps this same permissive nature for gene duplication and mutation has provided IFNT genes with heightened abilities to serve as pregnancy recognition factors. IFNT protein variants differ in antiviral and antiproliferative activities (Martal et al. 1990, Fillion et al. 1991, Ott et al. 1991, Alexenko et al. 1999, Winkelman et al. 1999). Also, prostaglandin (PG) production in cultured bovine endometrial cells is regulated to different extents by different IFNT variants (Parent et al. 2003). Most of the bovine and ovine IFNT variants studied thus far were proficient at reducing PGE and PGF2α concentrations when added to cultures at low concentrations, but preferentially stimulated PGE2 production at higher concentrations. One boIFNT variant inhibited PGE2 production at all concentrations while another stimulated both PGE and PGF2α at high concentrations. There likely are several other key physiological events that could be manipulated differently by different IFNT variants. IFNT-responsive genes in the ovine uterus are involved in controlling apoptosis, cellular growth, angiogenesis, vasculogenesis, inflammation and several other processes (Chen et al. 2006, 2007, Gray et al. 2006). There is no information available on how specific IFNT variants might control the signaling networks that operate these diverse biological processes.

There is evidence that IFNT variants differ in their antiluteolytic potency. An adequate *in vitro* model for antiluteolytic activity has not been developed.
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Table 1 Minimal effective amounts of Ovis aries IFNT and IFNA recombinant proteins at extending estrous cycle lengths in cyclic ewes.

<table>
<thead>
<tr>
<th>Protein name**</th>
<th>Protein (μg/day)</th>
<th>Antiviral Activity (IU/day)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IFNT4 (p3)</td>
<td>10</td>
<td>0.7x10^7</td>
</tr>
<tr>
<td>IFNT2c (p8)</td>
<td>50</td>
<td>1.7x10^7</td>
</tr>
<tr>
<td>IFNT1a (OTP-1)</td>
<td>100</td>
<td>1.0x10^7</td>
</tr>
<tr>
<td>IFNT2c (trophoblastin)</td>
<td>340</td>
<td>1.9x10^7</td>
</tr>
<tr>
<td>IFNT6c (p6)</td>
<td>300</td>
<td>4.3x10^7</td>
</tr>
<tr>
<td>IFNT11 (s4)</td>
<td>1000</td>
<td>11.8x10^7</td>
</tr>
<tr>
<td>IFNA1</td>
<td>175</td>
<td>0.7x10^7</td>
</tr>
</tbody>
</table>

*Data were taken from various sources from work completed by the same laboratory over several years (Ealy et al. 1998a,b, Martal et al. 1990, Ott et al. 1993, Winkelman et al. 1999, Green et al. 2005);

**Nomenclature used for naming these IFNs is provided in (Alexenko et al. 2000). GenBank accession numbers and associated sequences in Fig. 3 are as follows. IFNT4 (X56341.1; sheep 7), IFNT2c (X07920.1 for p8; M26386 for trophoblastin; sheep 10), IFNT1a (Y00287 and NM-001123399; sheep 9), IFNT6c (X56343.1; sheep 22), IFNT11 (M73241.1; sheep 1), IFNA (AY802984; NA);

***The uterine horn ipsilateral to a functional corpus luteum was cannulated. The cannula was exteriorized at the flank. Twice daily injections, each containing one-half the daily amount of IFN, was given. The tubing was flushed with carrier solution to ensure complete delivery of the protein.

so that studies defining the potency of IFNT variants have utilized an animal model that involves surgical placement of uterine cannulas into cyclic ewes and daily IFNT injections during diestrus. The maintenance of elevated circulating progesterone concentrations is used as the indicator of how long CL lifespan and subsequent estrous cycle length were extended. Table 1 shows the minimal effective amounts of IFNT needed to extend luteal function in several independent studies (Ealy et al. 1998a,b, Martal et al. 1990, Ott et al. 1993, Winkelman et al. 1999, Green et al. 2005). When examining protein potencies based on the amount of IFNT protein administered, the IFNT4 variant was clearly more effective than other IFNT proteins at extending CL function. A related variant termed IFNT2c, which differs in only 3 AAs from IFNT4, required 5- to 34-times more protein to achieve the same effect, depending on the recombinant protein used. A more distinct variant, termed IFNT6c, required 30-times more protein to achieve this effect. A fourth IFNT protein, termed IFNT11 (formerly known as IFNT-s4/p4) was not a potent antiluteolytic factor. This protein was studied because recombinant protein could be produced in great quantities from reconstituted bacterial inclusion bodies (i.e. insoluble protein pellet) (Li & Roberts 1994b, Niswender et al. 1997). Although the 5’ UTR of IFNT11 resembles that of other IFNT, its coding region contains similarity with both IFNT and IFNW (Leaman et al. 1992, Walker & Roberts 2009). It is also lowly expressed in ovine conceptuses, likely because it lacks a functional Ets2 site (Ezashi et al. 1998). Interestingly, despite its lack of effectiveness in extending estrous cycle length, IFNT11 is a potent antiviral and antiproliferative agent (Ealy et al. 1998b, Alexenko et al. 1999). Therefore, whatever mutations have caused it to lose its antiluteolytic potential has not interfered with its ability to function as an immune-regulatory factor. A recombinant ovine IFNA protein also contained antiluteolytic activity, but more of it was needed to produce an antiluteolytic response than with IFNT4 and IFNT2c (Table 1) (Green et al. 2005).

There has been some debate as to whether antiviral activity can be used as a guide to investigate antiluteolytic potential. Complicating this issue is that the purity, stability to freeze thawing (not recommended) and biological shelf lives of recombinant IFN proteins may vary from batch to batch and protein to protein. Accordingly, all IFNs should probably be assessed for antiviral activity before beginning an experiment. However, antiviral activity is not associated with antiluteolytic activity. The antiviral potency of type I IFNs also has very little merit on how it will perform as an antiproliferative or immunomodulatory factor (Ealy et al. 1998b, Winkelman et al. 1999).

It is noteworthy to also mention there is a good deal of biological variation in individual ewe responses in regard to the length of time before ewes return to estrus after an IFNT treatment (Ealy et al. 1998a,b, Winkelman et al. 1999, Green et al. 2005). Most IFNT treatment groups contain ewes that (1) do not respond, (2) have a modest response (<21 day interestrous interval) or (3) have a good response to the treatment (21 to >40 day interestrous interval) (Green et al. 2005). Usually 5–15% of treated ewes fail to respond to an otherwise adequate IFNT treatment. These ewes may represent ewes that would undergo early conceptus loss because they are incapable of responding to IFNT. The proportion of modest and good responders changes based on the type and dose of IFN treatment given. For example, a greater proportion of ewes exhibited >21 day interestrous intervals when treated with ovIFNT4 (10μg/day) than when treated with ovIFNA (175μg/day) (67% vs 25% good responders respectively) (Green et al. 2005). Also, there appears to be an association between the length of IFNT treatment and the length of extended CL function. Ewes provided IFNT until day 18 or 25 post-estrus were more likely to exhibit estrous cycle extension that lasted ≥40 days (Spencer et al. 1999, Green et al. 2005) than ewes only treated up to day 15 or 16 post-estrus (Ealy et al. 1998b, Winkelman et al. 1999). One explanation for this phenomenon is that there may be two important facets to IFNT action during early pregnancy. The first includes the initial rescue of the CL and the second involves other uterine- and/or ovarian-dependent actions that maintain CL function for several weeks. Perhaps the comprehensive uterine and ovarian responses to IFNT enable this second set of actions to prolong the period of pseudopregnancy observed in ewes.
Concluding remarks

The intention of this review has been to provide a sufficient perspective on the vast amount of knowledge gained in the past 30 years about the evolution of IFNT. The crucial step that created the primordial gene encoding IFNT probably occurred when a trophoblast-specific promoter/enhancer was inserted into the gene control regions of an ancestral IFNW. Subsequent point mutations, duplications and conversion events have led to the present complement of contemporary IFNT genes, which appear to comprise as few as three but as many as seven in cattle. Cattle, sheep and goats also contain numerous polymorphic forms of IFNT. There is compelling evidence that these different forms do not have identical, biological activities, including their abilities to extend estrous cycle length. Several questions about the need for so many IFNT variants still remain unanswered. Among them is characterizing the full extent of individual and combined IFNT protein actions in the uterus.

Supplementary data

This is linked to the online version of the paper at http://dx.doi.org/10.1530/REP-17-0292.

Declaration of interest

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