Lamarck and Immunity : Somatic and Germline Evolution of Antibody Genes

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Abstract

Current work on the mechanism of hypermutation of somatically rearranged antibody variable (V) genes shows that the most likely mechanism involves both direct DNA modification (deamination of cytosines to uracils by AID deaminase) and strand nicking plus mRNA editing (deamination of adenosine to inosine via the ADAR1 deaminase) coupled to a reverse transcription process to fix RNA sequence modifications in V gene DNA – most likely involving the repair enzyme DNA polymerase eta (η) known to be an efficient reverse transcriptase *in vitro*. The DNA sequence patterns of families of similar germline V genes reveals that many features of somatically mutated and antigen-selected variable genes appear written into the germline V gene arrays of the immune system. Lamarckian gene feedback and cellular reverse transcription, coupled to Darwinian antigen binding selection of somatically mutated V genes, are concepts which appear necessary for a more complete understanding how the V gene complex has evolved. Antibody variable (V) genes of the immune system have therefore been used to test ideas on reverse transcriptase-coupled soma-to-germline feedback in a complex multicellular system. Such feedback constitutes a violation of Weismann's Barrier and thus support for some type of Lamarckian gene feedback operative during the evolution of the vertebrate immune system.

Keywords: Lamarck, immunity, somatic and germline evolution, antibody genes; evolution

Introduction

In this paper I will review the main findings of our studies on the mechanism of antigen-driven somatic hypermutation of rearranged antibody (immunoglobulin, Ig) variable region genes (so called VDJs) and the impact of this somatic genetic diversity on the germline V segment repertoire. I will then draw general conclusions on the origins of genome diversity. Many details of this work have already been covered in major reviews (Steele *et al.* 1993; Rothenfluh *et al.* 1995; Steele *et al.* 1997; Blanden *et al.* 1998) and in our 1998 book *Lamarck's Signature* (Steele *et al.* 1998). More recent work will be cited in the body of the text.

General Concepts: Molecular-Cellular Immunology and Evolution

It is generally agreed that the primary evolutionary purpose of the immune system of vertebrates is the protection of the individual against disease. The proteins and carbohydrates which make up the cell walls, viral coats and secreted microbial toxins constitute the foreign antigens which individual immune systems need to react against to preserve the integrity of the body. The system consists of highly mobile blood white cells (lymphocytes) which come in two main categories, B cells and T cells which circulate system has been shaped by two selective forces, a) the requirement to respond to unexpected antigens thrown up by new infectious diseases, and b) the need to prevent autoimmune reactions against self antigens. This has meant that during vertebrate evolution the immune system has developed strategies to learn to recognise and respond to the antigenic universe both during ontogeny (somatic recognition strategy during life in individual animals) and phylogeny (a germ line strategy for antigen recognition). The founding concepts of modern immunology are based on the Clonal Selection Theory of Acquired Immunity of Sir MacFarlane Burnet (Burnet 1959). A good summary of the theory can be found in the *Scientific American* article by Ada and Nossal published in 1987.

from blood to lymph via a complex network of lymphatic vessels and capillaries. The complexity of the

system almost rivals that of the brain and central

nervous system (which in contrast consists of sessile or

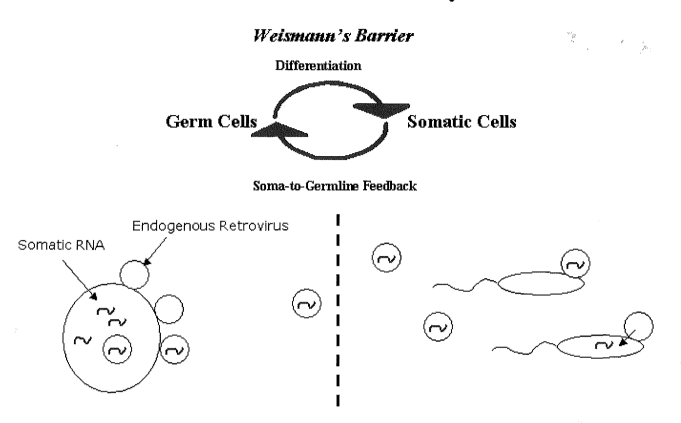
non-mobile nerve cells and fibres which generate their

complexity in both their sheer cell numbers and cell-cell

^{v synaptic connections). The progenitors of the white cell lineages (B and T lymphocytes, monocytes, neutrophils, mast cells, polymorphonuclear leucocytes} *etc.*) arise from stem cells in the bone marrow which produce many millions of hemopoietic cells on a daily basis (and, of course, the senescence of many other hemopoietic-derived cells as they exit the system). Hormonal and cytokine cell-to-cell communication no doubt allows the system to be co-ordinately controlled.
The primary evolutionary strategy of the immune system has been shaped by two selective forces, a) the requirement to respond to unexpected antigens thrown

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Weismann's Barrier Selectively Permeable



Somatic Selection Hypothesis, 1979

Figure 4. Somatic selection hypothesis, 1979.

gene arrays at the immunoglobulin locus in vertebrates. This work has allowed us to conclude that a significant portion of the somatic mutation and antigen-selection pattern in antibody variable genes (~80%) is indeed written into the germline V gene arrays at Ig loci.

Rearrangement, Gene Expression and Somatic Hypermutation of VDJ genes

A critical factor in the analysis depends on what has been established about the germline and somatic expression of immunoglobulin genes – they have clear 'germline' and 'somatic' configurations (Figure 5 and see Honjo *et al.* 2004). These facts allow us to infer and deduce that genetic information has indeed flowed from the somatic compartment to the germ cell compartment over evolutionary time.

The figure shows a schematic outline of a mammalian immunoglobulin heavy chain in its germline configuration and its somatic configuration. The germline, or unrearranged DNA configuration, exists in germ cells and all non-lymphoid cells in the body (*e.g.* kidney cells, liver cells, *etc.*, Honjo *et al.* 2004). Thus on the left hand side (5' side) are the array of so-called 'V-elements' or 'V-segments', which would typically encode approximately 95 amino acids and the 100–200 V-

elements are encoded in a span of chromosomal DNA of about 1 Mb in the human genome . This repertoire of unrearranged V-elements lies about 100Kb upstream (in the transcriptional sense) of very short genetic elements termed diversity (D) and joining (J) regions. There are 10-30 D regions and 4-5 J regions at typical mammalian IgH loci (together they would encode after VDJ assembly approximately 25 additional amino acids). Further downstream, encompassing about 10 Kb lie successively the intronic enhancer and nuclear Matrix Attachment Region (EiMAR) and then the Ig class switch region DNA repeat elements and then a series of constant region exons encompassing Ig heavy chain isotypes, mu (v), delta (δ), the various γ chain subclasses (IgG1, IgG2a, IgG2b IgG3) and then the α chains for secretory IgA and ε chains for mast cell binding and allergy-activating IgE antibodies. Further downstream is the 3' enhancer region (Honjo et al. 2004).

The various Ig classes reflect the functional properties of the antibody once antigen has been bound by the antigen combining site. The functional properties would include Complement activation and thus opsonisation of foreign particles for phagocytosis by monocyte scavenger cells (*e.g.* macrophages). The antigen binding site is a heterodimer of a light (L) and heavy (H) chain so antigen binding and thus antigen-mediated selection can only

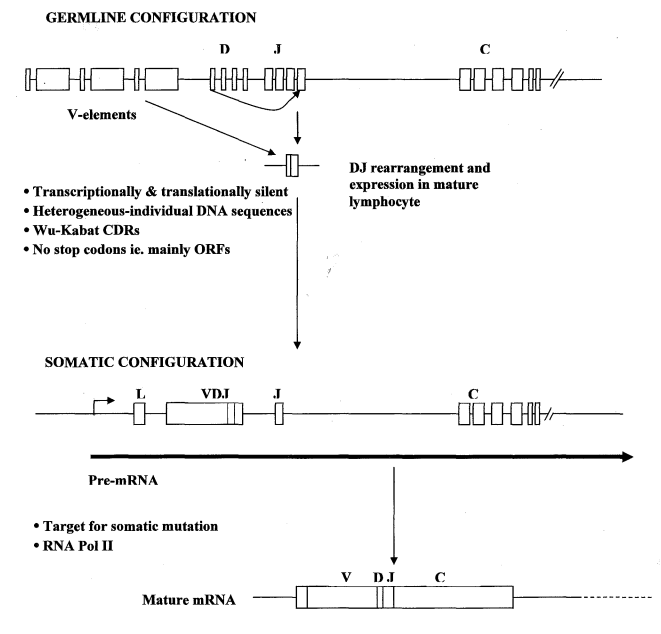


Figure 5. Rearrangement and immunoglobulin gene expression.

occur on a fully assembled Ig molecule or a B cell that displays such an antigen receptor on its surface membrane.

A key process not shown in Figure 5 is immunoglobulin class switching (CSR) whereby following cytokine signalling from other lymphocytes (T cells) and white cells, the B cell will switch from transcribing and assembling IgM heavy chains (υ chains), and reposition the productively assembled VDJ gene further downstream in front of one of the down stream Ig isotypes. This is a looping out DNA recombination mechanism such that the B cell retains the original selected VDJ but now has it joined to a different set of constant region exons (Honjo *et al.* 2004).

It is already clear that Ig loci display a degree of genetic complexity not observed in more straight forward single-copy house keeping or tissue-specific protein coding genes. T-cell receptor genes also display the same general genetic organisation and expression strategy (but typically do not normally somatically hypermutate their assembled VDJ genes). There is evidence that the protocadherin synaptic receptor genes in the central nervous system show a similar variable-to-constant rearranging strategy as seen for immunoglobulins and Tcell receptor genes but in these cases it is executed at the RNA level by an alternative splicing mechanism (for a mini-review see Chess 2005).

Germline V-elements *per se* are never transcribed into RNA for inclusion in a mRNA prior to translation into Ig proteins. In this sense they are transcriptionally and translationally silent. As such V-elements or their products are never the direct targets of antigen-binding selection. This type of antigen-mediated somatic selection is only directed to a fully assembled VDJ gene in the context of a light chain VJ gene co-expressed in the same B cell (and thus clonally selected).

VDJs are therefore the substrates for both RNA polymerase II transcription and somatic hypermutation. It is important to note that additional mutational errors are introduced by the DNA rearrangement process at the V-D and D-J borders (termed 'junctional diversity') and because the process is stochastic only a minor portion (about 10% of all rearrangements of IgH chains) are 'productive' *i.e.* in the correct translational reading frame (Honjo *et al.* 2004). This critical point will be discussed later in the context of fused 'VD' pseudogenes at chicken IgH loci (Rothenfluh *et al.* 1995).

Thus a germline repertoire of 100 functional VH elements, 20 D and 5 J regions can theoretically encode 100 x 20 x 5 or 10,000 VDJ regions; similarly 100 functional VL elements and 5 J regions would encode 500 VL. Together there is a potential combinatorial germline repertoire of perhaps 10,000 x 500 or 5 x 10⁶ unique antibody specificities. Following antigenic stimulation and somatic hypermutation this potential repertoire could perhaps increase another order of magnitude or two (Berek & Milstein 1987,1988). In reality ongoing antigen selection sifts and focuses the response for higher affinity antibodies so the full potential is unlikely to be realised in any individual (and of course the upper limit would be set by the total number of B cells generated from the bone marrow at any given point in time).

To summarise, genetic information in the form of unrearranged V-elements is never subject to direct antigen binding selection on the intact antibody or Ig receptor bearing cell. In contrast B cells expressing fully rearranged VDJ (heavy chain) and VJ (light chain) genes are subject to direct antigen-binding selection. It is this crucial distinction that demarcates the germline from the somatic configuration and thus allows deductions on the origin of highly non-random DNA sequence patterns.

The Germinal Centre and Affinity-Based Selection

A naïve B cell in the periphery can be selected by antigen to immediately secrete its encoded antibody or it can migrate to the primary follicle in lymphoid tissue to become a founder B cell in a Germinal Centre (termed GC). One or just a few B cells locate in a follicle and they multiply to form small colonies of 10,000 to 20,000 cells. Due to antigen-binding competition between pre-existing low affinity antibody and antigen-antibody complexes displayed on follicular dendritic cells within the GC, only the mutated B cells displaying viable high affinity antibodies survive - the rest die by the programmed cell death process called apoptosis (≥90% of all B cells in a Germinal Centre die there). In this way the mutated B cell survivors become antibody secreting cells and memory cells and they bear the signature of non-random DNA sequence modifications typical of intense selection. That is, point mutations in the VDJ accrue in those regions termed CDR or Complementary Determining Regions, which encode the amino acids that make direct contact with the molecular shapes of the antigen (typically protein, carbohydrate). Typical 'Wu-Kabat' plots of this non-random variability are shown in Figure 6 for 30 somatically mutated derivatives of the rearranged VH186.2 gene in mice (Steele et al. 1993). All of these features of the Germinal Centre reaction have

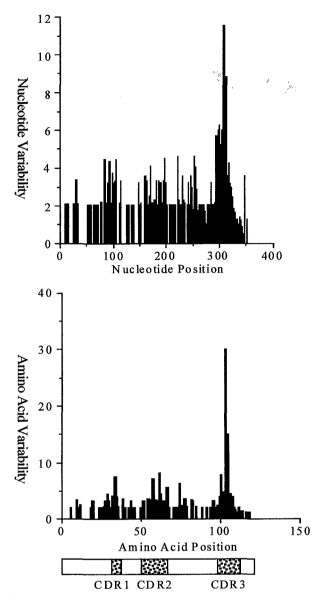


Figure 6. Wu-Kabat variability plots. Thirty somatically mutated derivatives of the mouse VH-1186.2 heavy chain variable gene assembled as V(D) is in mature anti-NP antibodies. Variability = Number difference at a position / Frequency of most common at that position. VDJ DNA sequences at top; translated protein sequences at bottom. Adapted from Steele *et al.* (1993).

been covered at length elsewhere eg. in *Lamarck's Signature* (Steele *et al.* 1998) or can be found in more specialised publications (*e.g.* MacLennan 1994).

The Mechanism of Somatic Hypermutation

The dominant current model of somatic hypermutation, "The DNA Deamination" model is DNAbased. *i.e.* all the mutational events occur directly at the DNA level (Di Noia & Neuberger 2007; Teng & Papvasiliou 2007). The main first step entails deamination of Cytosine to Uracil by the enzyme activation-induced cytidine deaminase (AID) which targets Cytosines in the context of WRCY hot spots (W = A or T, R = A or G purines, and Y = C or T pyrimidines). The resulting C-to-U lesions in DNA are either repaired by a base excision DNA repair pathway (involving uracil DNA glycosylase, UNG), or if not repaired, replicated over to produce C-to-T mutations. If repaired by UNG the resulting abasic site can be transformed into a nick in the DNA by an endonuclease termed apurinic apyrimidinic endonuclease (APE). Alternatively the G:U mispairs attract the mismatch repair heterodimer MSH2-MSH6 which also recruits the error-prone translesion DNA polymerase- η (eta) which introduces mutations in the repaired patch by targeting A:T base pairs at WAsites (where W = A or T). These series of steps are very similar to the V targeted-nicking and error-prone repair model of somatic mutation of IgV genes first advanced by Brenner and Milstein in 1966.

However three sets of recent observations by our group are not easily reconciled with the standard model but are consistent with, or predicted, by the reverse transcriptase model (Steele 2009):

- a) DNA polymerase-η the accepted and *sole* A:T mutator in SHM is an efficient reverse transcriptase *in vitro* (Franklin *et al.* 2004);
- b) The RNA editing signature of ADAR1-mediated A-to-I deamination, as instanced by the elevated A-to-G mutations, is embedded within the SHM pattern (Steele *et al.* 2006); and
- c) The AID deaminase-linked RNA polymerase II RNA mutation signature, as instanced by elevated mutations at G sites (particularly G-to-C and G-to-A) is embedded within the SHM pattern (Steele 2009).

This work has led us to conclude that the weight of evidence now favours "The Reverse Transcriptase Model first advanced by Steele and Pollard in 1987. Thus somatic hypermutation in B lymphocytes involves:

a) Direct DNA deamination (C-to-U, thus giving rise to C-to-T and G-to-A mutations);

- b) RNA Pol II copying deaminated DNA templates carrying U and abasic site lesions generates mutated mRNA (giving rise to G-to-C and G-to-A strand biased mutation signatures);
- c) RNA deamination (editing) of mRNA causing Adenosine-to-Inosine mutations via ADAR1 deaminase (thus causing the A-to-G strand biased mutation signature); and
- d) Error-prone reverse transcription by DNA Pol-η to fix the RNA mutations in B lymphocyte DNA and create further strand biased mutations *viz.* the transversions A-to-C and A-to-T.

To summarize, during somatic hypermutation both direct DNA mutations and a variety of RNA mutations are copied back into DNA (Steele 2009). As we have pointed out earlier (Steele *et al.* 1997) a role for a *cellular* reverse transcriptase such as DNA polymerase- η acting in its reverse transcriptase mode should no longer be a heretical concept given that "telomerase", a ribonucleic acid-protein particle has as its core function the capacity to copy the RNA repeat into a DNA repeat – a critical step in synthesis of the telomere multiple repeat chromosomal cap. That is "telomerase" itself is a cellular reverse transcriptase (Blackburn 1992).

Evolution of the Germline V-Segment Repertoire and Soma-to-Germline Feedback

Is there any evidence that somatically mutated variable genes can be fed back to the germline V-segment repertoire? It is possible to answer this question in the affirmative because germline V-segments can *never* be the targets of somatic hypermutation nor direct antigenmediated selection at the protein level *i.e.* on an intact Ig antigen receptor on a mature B cell surface. This fact is

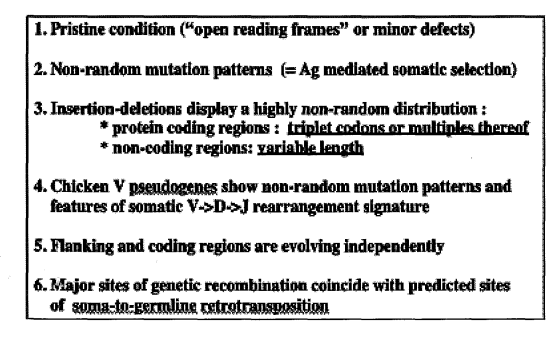


Figure 7. Key features of the repertoire Germline V Segments (Blanden et al. 1998).

often not addressed nor appreciated by those in the field speculating on the evolution of the V-segment repertoire. As we have documented elsewhere the germline V repertoires of families of similar V segments display all the hallmarks of strong somatic mutation and antigenmediated selection *i.e.* a significant portion of the somatically mutated VDJ repertoire generated during evolutionary time in vertebrates has been fed back to the germline most likely by a reverse transcriptase intermediate step and targeted to unrearranged V genes by homologous recombination (approx ≥80% of the assembled VDJ gene is comprised of the V element). Figure 7 summarises the main findings of this work which has been extensively published in refereed literature (Rothenfluh et al. 1994; Rothenfluh et al. 1995; Weiller et al. 1998; Blanden et al. 1998; Zylstra et al. 2003).

In short, a highly non-random somatic mutation and selection signature dominates the DNA sequence pattern of families of similar V genes arrayed, usually in tandem, in the vertebrate germline.

As suggested above these are 'subtle' somatic mutation signatures – they are of a different class from the more obvious retro-sequence impact events that dominate vertebrate and mammalian genomes. They argue for a requirement for innovative ways of interpreting the DNA landscape of the genome. Thus sites of hyperrecombination near RNA splice site borders in the L-V intron make sense in this model (Weiller *et al.* 1998) as do the strange features of chicken VH pseudogenes which all have fused 'D' bits in the correct reading frame (Rothenfluh *et al.* 1995; Ota & Nei 1995).

From our research thus far we therefore conclude:

- 1. Somatic hypermutation of antibody V genes operates by direct DNA and RNA base modifications coupled to reverse transcription and integration of mutated cDNA retrotranscripts back into chromosomal DNA *within* a B lymphocyte.
- 2. Over evolutionary time somatically mutated and selected ("successful") V sequences from B lymphocytes have undergone homologous recombination into germline DNA, thus contributing to germline diversity *and* the maintenance of a *functiona*l germline V gene repertoire.

Evidence from other systems

Is Soma-to-Germline feedback a general phenomenon in complex biological systems? We address this by briefly reviewing the work of other groups.

Corrado Spadafora

From about the late 1980s to the present Corrado Spadafora and colleagues in Rome have published a series of important papers clearly showing that mammalian spermatozoa can take up foreign nucleic acid molecules and express the genetic information in progeny organisms. In particular mouse spermatozoa *in vitro* can absorb both foreign DNA and RNA, and if the latter then a LINE-1-derived reverse transcription step will be executed copying the RNA into DNA. In a small number of cases ($\leq 10\%$) the DNA sequences are

integrated into the germline genome. In the majority of cases the sperm-absorbed DNA/RNA exists as extrachromosomal episomes which replicate along with the host somatic cells during development displaying mosaic tissue expression (see reviews in Smith & Spadafora 2005; Spadafora 2008). This work clearly shows there is no physical barrier to uptake of DNA or RNA, although there maybe clear developmental stages in spermatogenesis when spermatozoa are susceptible to foreign nucleic acid uptake (Zoraqi & Spadafora 1997).

Patrick Fogarty

Using an innovative technique based on P-elements and delivering DNA transgenes intravenously in simple vesicles, Fogarty has shown that 50% of progeny from such male mice inherit the gene sequence (Fogarty 2002). The critical integration event requires a transposase. This works suggests that non-cellular DNA can readily transverse the testes tissue barriers, that normally quarantine the production of sperm, be integrated into the germline and be transmitted to progeny.

Minoo Rassoulzadegan

The group of Minoo Rassoulzadegan has shown that mature sperm carry more than just a compact haploid DNA nucleus (Rassoulzadegan et al. 2006). Thus sperm heads contain gene-specific regulatory RNAs (miRNA) which at fertilisation can have profound genetic effects in progeny. The phenomenon described involves an allelespecific "paramutation" effect of the Kit locus (important in mammalian development) yet the implications of the finding are far reaching. Certainly the effects can be transmitted to an additional breeding generation. The mechanism is unclear given that animal miRNA systems are not thought to amplify their miRNA precursors by a double stranded RNA polymerase as in plants. Perhaps the transgenerational effects are based on long lived RNA molecules? However the whole phenomenon raises the possibility of germline fixation of such epigenetic intermediary effects via a reverse transcription step at some level of the RNA regulatory process.

John Mattick

Since the mid 1990s John Mattick and colleagues have been documenting the extent and importance of the RNA regulatory networks of what we now call the extended 'transcriptome" (Mattick 2007; Mattick et al. 2009). Thus non-protein coding regions produce ncRNAs which are regulatory in nature, regulating gene-specific expression of protein coding genes. Only about 2% of the entire mammalian 'transcriptome' codes for proteins - the rest (>98% genome) are involved in specific gene regulation in a multilayered complex best described as the "RNA regulatory universe". In more recent papers Mattick concedes the necessity for some form of soma-to-germline feedback to be operative (Mattick 2009) to ensure that selected genetic changes at this level contribute to the evolution of complex systems, particularly the brain and central nervous system (Mattick & Mehler 2008).

Lars Holmgren

For a number of years Holmgren and colleagues have studied the potential genetic consequences in metastases of horizontal transfer of tumor genes via dispersal and uptake of apoptotic bodies (Bersmedh *et al.* 2001; Ehnfors *et al.* 2009). These studies clearly show that if the laterally spreading DNA confers a selective advantage on the recipient cell then integration of the DNA is manifest and the DNA sequences propagated to progeny cells. We speculated on this type of somatic gene transfer in the late 1970s (Steele 1979).

Genetic Cargo of Sperm?

This is an appropriate question given the interesting findings of Spadafora and Rassoulzadegan we have just discussed. Indeed when one considers an ovum at about the time of natural fertilization there are more questions than answers raised by the phenomenon (Figure 8).

Have we become so used to an image like this that we have forgotten how truly amazing it really is. Apparently one sperm succeeds in the race to fertilisation (one possibility is arrowed). Given conventional wisdom all the other attached sperm have no further say in the genetic outcome. Apart from the non-Mendelian routes of genetic transfer in the experiments of Spadafora and Rassoulzadegan we have known for many years (e.g. Keissling et al. 1987) that sperm heads have clusters of attached endogenous retroviruses - to what end one might ask? Moreover, ERV concentrations are very high in seminal fluid (≥10¹¹ per ml) and ERVs are emitted in copious quantities from activated lymphocytes (they also are prominent in Germinal Centres following immunisation). ERVs have been observed coating the female placenta (see Rothenfluh 1995 for more references of this type). Again the question arises - to what biological purpose should cells of the immune system and reproductive tissue be so predominantly associated with either ERV production or unexpected ERV tissue localisation?

Concluding Remarks

We conclude that both Darwinian antigen-binding selection and Lamarckian soma-to-germline feedback play key roles in the evolution of antibody variable

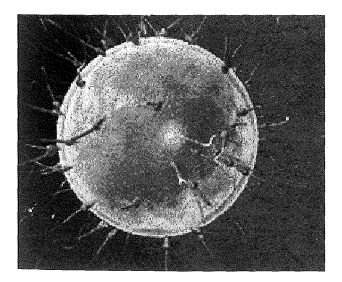


Figure 8. Sperm fertilising ovum.

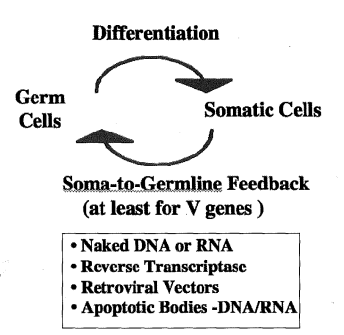


Figure 9. Weismann's Barrier selectively permeable.

genes. There is also evidence supporting the view that reverse transcription is central to a better understanding of the somatic and germline evolution of these genes. The work of a number of groups suggest that the ease of gene movement between cells, whether they be germline or somatic, suggests that soma-to-germline feedback is likely to be general in complex biological systems and contributes to genome diversity. Thus Weismann's Barrier is viewed as being *selectively permeable* to somatic genetic information provided it is beneficial to both parent and progeny organisms (Figure 9). Acquired somatic genetic information ("experience") may therefore not be lost with the death of the individual but be propagated to progeny who would then be selected in a Darwinian manner for fitness.

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