## Reverse Transcriptase Mechanism of Somatic Hypermutation: Sixty years of Clonal Selection Theory

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**Abstract:** The evidence for the reverse transcriptase mechanism of somatic hypermutation is substantial. In this 60th anniversary year of the publication of Burnet's Clonal Selection Theory the evidence is briefly reviewed and updated.

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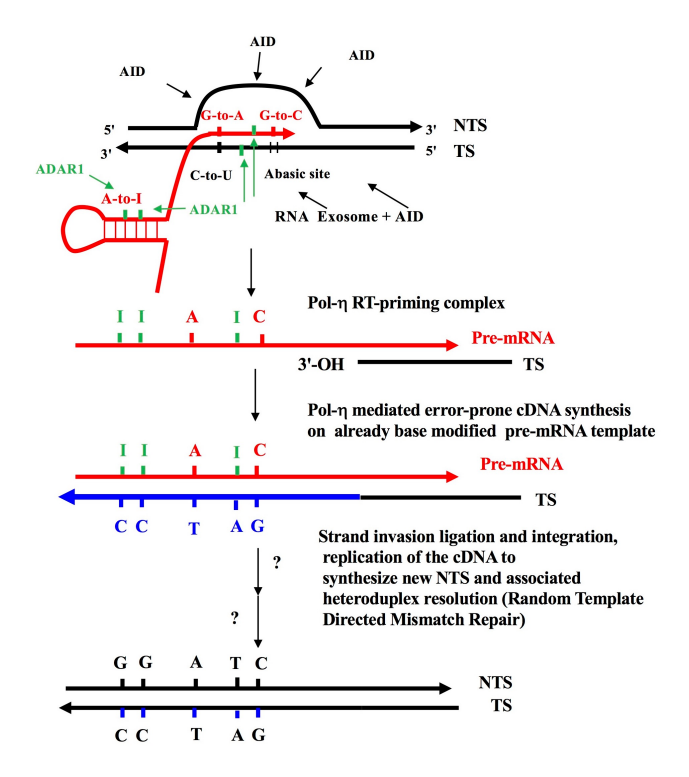
## Abbreviations used in this paper:

A>>T, mutations of A exceed mutations of T, e.g. as found in normal murine physiological SHM in vivo by 2.9 fold; ADAR, Adenosine Deaminase that acts on RNA; AID, activation induced cytidine deaminase, a APOBEC family member, initiating via C-to-U lesions at WRCY/RGYW Csite motifs in ssDNA of class switch recombination (CSR) and somatic hypermutation (SHM) processes at somatically rearranged Ig V(D)J gene loci; APOBEC family, generic abbreviation for the deoxyribonucleic acid, or dC-to-dU, deaminase family of which AID is a member (APOBEC3 A, B, C, D, F, G, H) similar in DNA sequence to the "apolipoprotein B RNA editor" APOBEC1; AP, an Abasic, or apurinic/apyrimidinic, site; APE, AP endonuclease; A-to-I, adenosine-to-inosine RNA editing; BCR, B cell Ig receptor; C, exons encoding constant regions of Ig molecules; D, the small "diversity' element part of the VDJ rearrangement process; G>>C, mutations of G exceed mutations of C, e.g. as found in normal murine physiological SHM in vivo by 1.7 fold; I, Inosine; Ig, immunoglobulin; IGHV, IGLV, megabase length germline regions encoding unrearranged V element arrays and associated D,J, C regions; J, joining elements, part of the VDJ rearrangement process; MMR, mismatch repair; MSH2-MSH6, MutSα heterodimer recognizing mispaired bases in DNA duplex; NTS, the non-transcribed, or "Top", 5' to 3' strand; Pol $-\eta$  or DNA polymeraseη (eta); RNA Pol II, RNA Polymerase II; R, Adenosine or Guanine; RT, reverse transcriptase; RT-Pol-n, reverse transcriptase activity displayed by Pol-n; SHM, somatic hypermutation; TS, the transcribed, or "Bottom", 3' to 5' strand, in context of a Transcription Bubble; TSRT, target site reverse transcription; U, uracil; UNG, uracyl DNA glycosylase involved in Base Excision Repair at dU sites in DNA resulting in either an Abasic site (AP) or APE-mediated ssDNA nicks (above); V[D]J, generic symbol for a rearranged immunoglobulin (or T cell receptor, TCR) variable region gene in the Adaptive Immune System; W, weak base pair involving A or U/T; WA-site, target motif for ADAR deaminase including DNA Polymerase-eta error prone incorporation in vitro; Y, pyrimidines T/U or C.

It is now 60 years since Sir MacFarlane Burnet first published *The Clonal Selection Theory of* Acquired Immunity, the foundation stone of modern immunology. Somatic mutation of the immunoglobulin variable region genes has been part and parcel of his clonal selection concept since its inception and is central to a rational understanding of immunological diversification, selftolerance and the emergence of cancer. We now have a very good idea of the molecular mechanism of somatic hypermutation. I have chosen to fit this scientific progress within 60 key publications since the late 1950s (Table 1). The most likely and plausible central molecular mechanism of Ig SHM, that fits with and explains all the evidence <sup>58</sup> is based on "Reverse Transcription" of the base-modified Ig pre-mRNA (Figure 1). That is, error-prone reverse transcription, by DNA Polymerase- $\eta$ , of the Ig pre-mRNA template intermediate at rearranged V(D)J gene somatic loci. The Ig pre-mRNA encoding the V(D)J region is copied off the transcribed DNA strand carrying prior AID C-to-U deamination lesions (Uracils and Abasic sites), and it also accumulates ADARdeaminase mediated RNA editing A-to-I modifications. This already base-modified pre-mRNA sequence is then copied back to the B lymphocyte genomic DNA and integrated at the rearranged VDJ site (concurrent with antigen-mediated selection of BCR bearing B lymphocytes, Centrocytes, in the Germinal Center). This is essentially the "Reverse Transcriptase Mechanism" which Jeff Pollard and I first published 30 years ago. <sup>21</sup> The mechanistic steps, many logical, are clearly outlined in Figure 1 which shows that the A>>T and G>>C strand bias-generating mutagenic activity is firmly focused on the nascent RNA intermediate in the context of the Transcription Bubble. 52,52,54,58,60 Recent publications should be consulted for further recent definitive ADAR Ato-I editing of RNA and DNA moieties at RNA:DNA hybrids within Transcription Bubbles. 58-60 Not only is it important to understand the correct molecular mechanism of SHM for cancer diagnosis and detection <sup>57,64</sup> but also to the current efforts to better understand <sup>31,65</sup> the origin of Ig diversity involving the mechanism of evolution of the sets germline V segments and the long IGHV and IGLV haplotypes in individual human beings. <sup>66,67</sup>. The author welcomes discussion by email.

Table 1 History of Somatic Hypermutation: Developments Relevant to The Reverse Transcriptase Mechanism

| Year    | Author                       | Main Development-Discovery-Concept   | Ref.   |
|---------|------------------------------|--|--------|
| 1957-59 | FM Burnet                    | Somatic mutation concept for Ab mainly in context "forbidden" anti-self clones       | 1      |
| 1959    | J Lederberg                  | Somatic mutation explicit in lymphocyte development and Ab diversity                 | 2      |
| 1962    | JB Fleishman et al           | Amino acid variation in N -terminal regions of V or antigen binding regions          | 3      |
| 1966    | S Brenner, C Milstein        | Model: V region specific nicking and error prone repair - "somatic hypermutation"    | 4      |
| 1967    | O Smithies                   | Somatic "Master->Slave" Gene Recombination model Ab diversity                        | 5      |
| 1967    | GM Edeleman, JA Gally        | Somatic recombination between duplicated V gene model Ab diversity                   | 6      |
| 1968    | M Cohn                       | Molecular biology of expectation - rational for SHM and response to unexpected       | 7      |
| 1970    | MG Weigert et al             | Somatic variability in Lambda light chain V region protein sequences                 | 8      |
| 1970    | TT Wu, EA Kabat              | Hypervariable regions coincide with and define antigen contact regions               | 9      |
| 1974    | AJ Cunningham                | The generation of antibody diversity after antigen                                   | 10     |
| 1974    | M Cohn                       | Somatic mutation explanation for Ab diversity clearly laid out                       | 11     |
| 1976    | S Tonegawa, C Steinberg      | DNA V gene counting confirms somatic mutation at molecular level in V lambda         | 12     |
| 1977    | S Tonegawa et al             | DNA V gene counting confirms somatic mutation at molecular level in V lambda         | 13     |
| 1981    | PJ Gearhart et al            | Somatic hypermutation of the TEPC15 VH rearranged gene in vivo                       | 14     |
| 1981    | ALM Bothwell et al           | Somatic hypermutation to the VH186.2 VH rearranged gene in vivo                      | 15     |
| 1981    | E Selsing, U Storb           | Somatic hypermutation of the MOPC167 VK rearranged gene in vivo                      | 16     |
| 1982    | PJ Gearhart                  | Somatic Hypermutation in Rearranged (VDJ) Variable Region Genes In Vivo              | 17     |
| 1983    | PJ Gearhart,DF Bogenhagen    | Somatic mutations occur in the 5' and 3' non-ding regions around VDJ genes           | 18     |
| 1985    | C Berek, C Milstein          | Use of hybridoma technique to sample somatic V[D]J mutant generation in vivo         | 19     |
| 1986    | A Cumano, K Rajewsky         | Further use hybridoma technique to sample somatic VDJ mutants in vivo                | 20     |
| 1987    | EJ Steele, JW Pollard        | Model: The reverse transcriptase mechanism of somatic hypermutation                  | 21     |
| 1987    | Golding et al                | First hint of strand biases in somatic hypermutation patterns viz. A>G versus T>C    | 22     |
| 1990    | GW Both et al                | Defining the 5' and 3' boundaries of somatic hypermutation at VDJ genes              | 23     |
| 1990    | SG Lebecque, PJ Gearhart     | Defining 5' and 3' boundaries of somatic hypermutation at VDJ genes                  | 24     |
| 1991-96 | IB Rogozin et al             | Identification RGYW/WRCY and WA hotspots in somatic hypermutation data               | 25,26  |
| 1992    | EJ Steele et al              | Defining the asymmetrical 5' to 3' somatic mutation distribution around V[D]J genes  | 27     |
| 1993    | AG Betz et al                | Defining the mutational hot spots across mutated V[D]J transgenes genes              | 28     |
| 1995    | J Yelamos et al              | Any non-Ig sequences parked between Promotor and J-C intron somatically mutate       | 29     |
| 1996    | A Peters, U Storb            | Strong evidence that transcription of VDJ target regions allows somatic mutation     | 30     |
| 1995-98 | GF Weiller, RV Blanden et al | The SHM signature is written into the germline V segment array                       | 31     |
| 1998    | C Milstein et al             | Both DNA strands targeted for G:C and A:T mutations in somatic hypermutation         | 32     |
| 1998    | Y Fukita et al               | Strong correlative evidence that transcription of VDJ allows somatic mutation        | 33     |
| 1998    | C Rada et al                 | In MSH2-deficient mice mutations are G:C focused suggesting two stages SHM           | 34     |
| 1999    | C Masutani et al             | Discovery of DNA Polymerase -eta and Y family translesion polymerases                | 35     |
| 2000    | M Muramatsu et al            | AID discovered - required to intiate SHM and Ig Class Switch Recombination           | 36     |
| 2001-2  | IB Rogozin, Y Pavlov et al   | Error-prone DNA Polymerase eta SHM spectrum correlates with WA hotspots              | 37,38  |
| 2001    | X Zeng et al                 | DNA Polymerase eta is the A:T mutator in somatic hypermutation in humans             | 39     |
| 2002-4  | MS Neuberger et al           | Definitive evidence that AID is a direct DNA C-to-U deaminase of the APOBEC family   | 40     |
| 2003    | R Bransteitter et al         | AID deaminates C>U on ssDNA - targets displaced strand Transcription Bubble          | 41     |
| 2003    | J Chaudhuri et al            | AID deaminates C>U on ssDNA - targets displaced strand Transcription Bubble          | 42     |
| 2003    | SK Dickerson et al           | AID deaminates C>U on ssDNA - targets displaced strand Transcription Bubble          | 43     |
| 2004    | J Chaudhuri et al            | AID deaminates C>U on ssDNA - targets displaced strand Transcription Bubble          | 44     |
| 2004    | HM Shen, U Storb             | AID targets both strands at Transcription Bubbles during transcription VDJ           | 45     |
| 2004    | C Rada et al                 | MSH2-MSH6 -/-and Uracil DNA Glycosylase -/-define G:C and A:T mutation phases        | 46     |
| 2004    | A Franklin et al             | Human DNA Polymerase eta is an efficient reverse transcriptase, as are kapp, iota    | 47     |
| 2004    | EJ Steele et al              | First hint that A>G versus T>C strand bias involves an A>I RNA edited intermediate   | 48     |
| 2005    | TM Wilson et al              | MSH2-MSH6 stimulates DNA polymerase eta, suggesting a role for A:T mutations         | 49     |
| 2006    | EJ Steele, RA Lindley et al  | Evidence WA>WG mutations correlate with the number nascent WA RNA stem loops         | 50     |
| 2007    | F Delbos et al               | Evidence that DNA Polymerase eta is the sole error-prone A:T SHM mutator in vivo     | 51     |
| 2009    | EJ Steele                    | SHM data 1984-2008 shows A>>T, G>>C strand biases explained by RNA/RT-model          | 52     |
| 2010-13 | EJ Steele, RA Lindley        | A>>T, G>>T SHM strand biases evident in non-Ig genes across all cancer exomes        | 53, 54 |
| 2011    | U Basu et al                 | RNA exosome exposes ssDNA for AID on transcribed strand at Transcription Bubbles     | 55     |
| 2011    | RW Maul et al                | AID generated Uracils physically located in the DNA of VDJ & Ig class switch regions | 56     |
| 2013    | RA Lindley                   | Codon-context targeted somatic mutation (TSM) in cancer exomes                       | 57     |
| 2016    | EJ Steele                    | Extant evidence supports the RNA/RT-based model and not the DNA-based model          | 58     |
| 2017    | Zheng et al                  | ADAR can directly edit both RNA and DNA A-sites in RNA:DNA hybrids                   | 59     |
| 2017    | EJ Steele, RA Lindley        | ADAR A>I Editing at RNA:DNA Hybrids is strong support for RNA/RT-based model         | 60     |
|         |                              |  |        |



**Figure 1 Legend - The Reverse Transcriptase Mechanism of Somatic Hypermutation**Modified in part from Figure 1 in Lindley and Steele.<sup>54</sup> This is an adaptation of the target site reverse transcription (TSRT) process of Luan et al. <sup>61</sup> Shown is a RNA Polymerase II generated Transcription Bubble with C and A substrate deamination by AID and ADAR proteins; and the sequelae showing some key hypothesized DNA and RNA intermediates highlighted for the generation of the main strand- biased mutation signatures involving A-to-G, G-to-A, G-to-T and G-to-C. <sup>52,58,60</sup> Black lines are DNA strands, red lines are pre-mRNA, blue thick lines are cDNA strands copied off pre-mRNA by reverse transcription via DNA polymerase η <sup>47</sup> Green bars are Inosines. Shown also is the action of the RNA exosome <sup>55</sup> allowing access of AID deaminase to

cytosines on the Transcribed Strand (TS). The ssDNA regions on the displaced Non-Transcribed Strand (NTS) are established targets of AID action. 41-44 With respect to the RNA intermediary step in this process mutations are first introduced at the DNA level by AID-mediated C-to-U deaminations, and then uracil DNA glycosylase (UNG)-generated Abasic sites in the TS (which can further mature into single strand nicks via the action of AP endonuclease generating the 3'-OH in the TS). These template Uracil and Abasic sites are transcribed into pre-mRNA by RNA Pol II generating G-to-A and G-to-C modifications respectively in the pre-mRNA as shown in Kuraoka et al. 62 which on TSRT-mediated reverse transcription, 61 integration and DNA replication result in G-to-A and G-to-C mutations in the NTS, in a strand biased manner. <sup>52,54,58,60</sup> Separately, adenosine-to-inosine (A-to-I) RNA editing events at WA targets, mediated by ADAR1 deaminase, in the nascent pre-mRNA emerging from Transcription Bubble-proximal dsRNA stem loops may be copied back into DNA by reverse transcription via Pol-η. 50 In theory ADARs can also deaminate the RNA and DNA moieties in the RNA: DNA hybrid. 59,60 The strand invasion and integration of newly synthesized cDNA TS, as well as random-template mismatch repair <sup>63</sup> are hypothesized additional steps (not shown here). In more detail: RNA Pol II introduces mutations in the Ig pre-mRNA as it copies the AID lesions in TS DNA, concurrently A-to-I RNA edited sites appear in RNA stem(-loops) forming in nascent pre-mRNA near the transcription bubble <sup>50</sup> as well as in RNA:DNA hybrids within the bubble. <sup>59,60</sup> Next, the RT-priming substrates are formed by annealing the nicked TS strand with an exposed 3'-OH end (for Y Family translesion DNA polymerase- $\eta$ , 35 now acting in it reverse transcriptase mode. 47 These could arise due to excisions at previous AID-mediated Abasic sites, or due to an excision introduced by endonuclease activity associated with the MSH2-MSH6 heterodimer engaging a U:G mispaired lesion. <sup>49</sup> This allows extension of a new TS by cDNA synthesis from the 3'-OH end copying the already base modified pre-mRNA template (with Inosine base pairing preferentially, like G, with C). Show is an A>T transversion generated at the RT step at a template Inosine.

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