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A plasma cell differentiation quality control ablates B cell clones with biallelic Ig rearrangements and truncated Ig production

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Aberrantly rearranged immunoglobulin (Ig) alleles are frequent. They are usually considered sterile and innocuous as a result of nonsense-mediated mRNA decay. However, alternative splicing can yield internally deleted proteins from such nonproductively V(D)J-rearranged loci. We show that nonsense codons from variable (V) Igκ exons promote exon-skipping and synthesis of V domain-less κ light chains (ΔV-κLCs). Unexpectedly, such ΔV-κLCs inhibit plasma cell (PC) differentiation. Accordingly, in wild-type mice, rearrangements encoding ΔV-κLCs are rare in PCs, but frequent in B cells. Likewise, enforcing expression of ΔV-κLCs impaired PC differentiation and antibody responses without disturbing germinal center reactions. In addition, PCs expressing ΔV-κLCs synthesize low levels of Ig and are mostly found among short-lived plasmablasts. ΔV-κLCs have intrinsic toxic effects in PCs unrelated to Ig assembly, but mediated by ER stress–associated apoptosis, making PCs producing ΔV-κLCs highly sensitive to proteasome inhibitors. Altogether, these findings demonstrate a quality control checkpoint blunting terminal PC differentiation by eliminating those cells expressing nonfunctionally rearranged Igκ alleles. This truncated Igκ exclusion (TIE) checkpoint ablates PC clones with ΔV-κLCs production and exacerbated ER stress response. The TIE checkpoint thus mediates selection of long-lived PCs with limited ER stress supporting high Ig secretion, but with a cost in terms of antigen-independent narrowing of the repertoire.

During early B cell maturation, Ig loci undergo programmed DNA rearrangements of germline V (variable), D (diversity), and J (joining) gene segments. This error-prone program generates random V(D)J junctions and a diversified primary antibody repertoire (Jung et al., 2006). Functional Ig heavy (H) and light (L) chains are controlled at the stages of pre–B cell receptor (preBCR) and BCR expression, respectively (Melchers et al., 2000). These early checkpoints ensure expansion of B cells expressing functional Ig chains, while limiting the development of autoreactive clones (Rajewsky, 1996). Once positively selected, immature B cells transit to the periphery and join the mature B cell pool. Upon antigen (Ag) stimulation, germinal center (GC) B cells diversify their receptors through activation-induced cytidine deaminase (AID)–dependent somatic hypermutation (SHM) and class-switch recombination (CSR; Manis et al., 2002; Pavri and Nussenzweig, 2011; Andrews et al., 2013). Self-reactive clones are also tightly controlled in mature and GC B cells, and BCR-signal strength regulates these late tolerance checkpoints, eventually leading to anergy or elimination of B cells (Allen et al., 2007; Victora and Nussenzweig, 2012). Our group has recently demonstrated that a recombination by IgH locus suicide recombination (LSR) participates actively in the late elimination of GC B cells (Péron et al., 2012). Cells that survive negative selection further differentiate into either memory B cells or plasma cells (PCs) secreting high-affinity antibodies. PCs are antibody-producing factories in which a massive expansion of the endoplasmic reticulum (ER), together with elevated production of chaperone proteins such as GRP78/BiP (glucose-regulated protein 78 kD/binding immunoglobulin protein), ensures proper folding and secretion of large amounts of Ig (Gass et al., 2002; Ron and Walter, 2007; Todd et al., 2009). Major transcriptional changes accompany PC differentiation, including a boost of Ig gene expression and modification of their splicing pattern, from membrane-type toward secretory-type Ig mRNAs (Santos et al., 2011).

Random nucleotide additions or deletions at V(D)J junctions inherently yield frameshifts and premature stop codons in two thirds of cases (Jung et al., 2006). When a nonproductive

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Abbreviations used: Ag, antigen; AID, activation-induced cytidine deaminase; AS, alternative splicing; CHOP, C/EBP homologous protein; CSR, class-switch recombination; ΔV-κLC, variable-domain-less κ light chain; ER, endoplasmic reticulum; GC, germinal center; κLC, κ light chain; LSR, locus suicide recombination; NAS, nonsense-associated altered splicing; NMD, nonsense-mediated mRNA decay; PC, plasma cell; PI, proteasome inhibitor; PTC, premature termination codon; SHM, somatic hypermutation; TIE, truncated Ig exclusion.

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V(D)J junction first affects one Ig allele, the second allele can still productively rearrange and support B cell maturation. Such biallelic V(D)J rearrangements, including a nonproductive allele, are retrieved in ∼40–50% of B lymphocytes (Mostoslavsky et al., 2004; Daly et al., 2007). It is now well accepted that nonproductive Ig alleles are actively transcribed during B cell development (Singh et al., 2003; Delpy et al., 2004a,b; Daly et al., 2007; Eberle et al., 2009; Tingueley et al., 2012; Holwerda et al., 2013). The nonsense-mediated mRNA decay (NMD) pathway ensures efficient degradation of the resulting Ig mRNAs harboring premature termination codons (PTCs), and hence limits the production of truncated Ig with C-terminal deletions (Li and Wilkinson, 1998). NMD likely protects lymphoid cells from adverse effects of aberrant Ig transcripts. Indeed, disruption of lymphocyte development was observed upon either expression of an UPF1 (up-frameshift protein 1) dominant-negative isoform (Frischmeyer-Guerrero et al., 2011) or conditional deletion of Upf2 (Weischenfeldt et al., 2008), two NMD likely protects lymphoid cells from adverse effects of aberrant Ig transcripts. Indeed, disruption of lymphocyte development was observed upon either expression of an UPF1 (up-frameshift protein 1) dominant-negative isoform (Frischmeyer-Guerrero et al., 2011) or conditional deletion of Upf2 (Weischenfeldt et al., 2008), two major NMD actors. In addition, Lutz et al. (2011) observed that the persistence of nonsense pI mRNA transcripts escaping effective NMD degradation impairs pro-B cell differentiation.

Abnormal RNA splicing is elicited upon recognition of nonsense mutations by the RNA surveillance machinery, leading to the accumulation of unspliced PTC-containing premRNAs or to the appearance of alternatively spliced mRNAs (Valentine, 1998; Maquat, 2002; Lejeune and Maquat, 2005). Several studies, including ours, have documented that in addition to NMD, nonproductive Ig transcripts are surveyed by the cooperative action of splicing inhibition and nonsense-associated altered splicing (NAS; Aoufouchi et al., 1996; Mühlemann et al., 2001; Bühler and Mühlemann, 2005; Chemin et al., 2010; Tingueley et al., 2012). Unlike NMD, which prevents the translation of prematurely terminated Ig polypeptides, nothing is known about the consequence of NAS and exon skipping with regard to the production of Ig mRNAs and proteins with internal deletions. Production of shortened Ig chains and exon skipping occurs in pathologies such as heavy chain diseases and monoclonal Ig deposition diseases (Cogné et al., 1988, 1992). In contrast, truncated Ig chains have never been documented in normal B cells, and whether aberrant Ig transcripts from excluded alleles end up with aberrant peptides is an open question. To address this issue during B cell development and PC differentiation, we analyzed the splicing pattern of Igκ transcripts. These simplest RNA molecules include only three exons: leader (L), variable (V), and constant (C). We found that the presence of nonsense codons within the V exon modified the splicing of Igκ transcripts and strongly enhanced exon skipping. Interestingly, the production of V domain-less κ light chains (∆V-κLCs) exacerbated the ER stress response and severely impaired PC differentiation. Thus, the transcription of nonproductively rearranged Ig alleles can be detrimental at some differentiation stages. Collectively, these findings reveal a late quality control process dedicated to truncated Ig exclusion (TIE). This TIE checkpoint blunts ∼20–25% of terminally differentiated PCs by eliminating in an Ag-independent manner half of those 40–50% cells with biallelic rearrangements of Igκ alleles.

RESULTS
Nonsense codons located within V regions enhance exon skipping of Igκ transcripts
We previously showed in normal B lineage cells that upon skipping of some noncoding V exons, so-called excluded alleles often yield shortened κ mRNAs which would translate into ∆V-κLCs (Chemin et al., 2010). Observing such shortened mRNAs is surprising because the production of truncated Ig is usually considered a specific feature of some lymphoproliferative diseases (Cogné et al., 1992). To extend this study, we further analyzed the conditions of exon skipping, checked whether shortened Igκ transcripts were translated into ∆V-κLCs, and addressed whether the latter affects B cell differentiation. Out-of-frame Vκ-to-Jκ rearrangements result in nonsense codons affecting either the 3′-end of the V exon (VPTC) or the C exon (CPTC; Delpy et al., 2004a; Fig. 1 A). We compared ∆V-κLC mRNA levels in cell lines transfected with constructs mimicking nonproductive VκJκ5 rearrangements from both the VPTC and the CPTC class (Fig. 1, B and C). This revealed high ∆V-κLC mRNAs in VPTC-expressing cells, whereas such transcripts were barely detectable in CPTC-expressing cells and absent in cell lines expressing productive Igκ genes (Fig. 1 D). ∆V-κLC mRNA levels were quantified using capillary electrophoresis after normalization to the amount of productive κLC mRNAs, as previously described (Chemin et al., 2010). Compared with CPTC-expressing cells, ∆V-κLC mRNAs were ∼3.5- and 6.7-fold higher in VPTC-expressing cells, whereas such transcripts were barely detectable in CPTC-expressing cells and absent in cell lines expressing productive Igκ genes (Fig. 1 D). ∆V-κLC mRNA levels were quantified using capillary electrophoresis after normalization to the amount of productive κLC mRNAs, as previously described (Chemin et al., 2010). Compared with CPTC-expressing cells, ∆V-κLC mRNAs were ∼3.5- and 6.7-fold higher in VPTC-expressing B (A20) and PC (S194) lines, respectively (Fig. 1 E). Hence, skipping of the V exon occurs mostly when this exon itself carries the PTC, and with an efficiency approximately two times higher in PCs than in B cells.

Exacerbated ER stress response in PCs expressing ∆V-κLCs
Next, we checked whether ∆V-κLC mRNAs were indeed translated into ofV domain-less truncated Ig, thus expected to mostly show up in VPTC-expressing PCs. Consistent with data in Fig. 1, complete κLCs (∼25 kD) and truncated ∆V-κLCs (∼12 kD) were detected in cells transfected with productive and VPTC constructs, respectively (Fig. 2 A). In contrast, no ∆V-κLCs (or a background level similar to nonsecretory Sp2/0 cells) was observed in untreated CPTC-expressing cells (Fig. 2 A). We further explored whether ∆V-κLCs were routed toward proteasomal degradation. In VPTC-expressing cells, treatment with MG132 induced an approximately two-to four-fold increase of ∆V-κLCs in cell pellets and culture supernatants, respectively (Fig. 2, A and B). In contrast in CPTC-expressing cells, very low amounts of ∆V-κLCs were found in cell pellets, and only upon treatment with MG132 (Fig. 2 A). For controls, no major changes were observed for complete κLCs (Fig. 2, A and B).

The survival of PCs depends on their ability to alleviate the ER stress response associated with massive Ig synthesis...
(Cenci et al., 2011). PC survival might thus be impacted by the production of proteins with abnormal structures and folding. To address this issue, we analyzed the apoptotic index of S194 PCs with regard to the abundance of ΔV-κLCs. Interestingly, VPTC-expressing cells exhibit higher apoptosis than nontransfected and CPTC-expressing cells, upon treatment with MG132 (Fig. 2 C). In addition, mRNA levels of C/EBP homologous protein (Chop), a transcription factor involved in ER-stress induced apoptosis (Obeng et al., 2006), were approximately two-fold higher in VPTC-expressing cells than in nontransfected cells upon treatment with MG132 (Fig. 2 D).

As expected, Chop expression in CPTC-expressing cells was intermediate, in between nontransfected and VPTC-expressing cells (Fig. 2 D). Therefore, the presence of ΔV-κLCs likely provokes excessive ER-stress that sensitizes plasma cells to CHOP-dependent apoptosis upon proteasome inhibition.

A TIE checkpoint counterselects those PCs expressing VPTC-rearranged Igκ alleles
Differential sensitivity to apoptosis can hardly be followed in vivo due to immediate phagocytosis of apoptotic cells. However, such differences would be expected to result in a bias in

**Figure 1.** Alternative splicing of nonproductive Igκ transcripts in B cell and plasma cell lines. (A, left) Mouse Igκ locus showing Vκ segments (in normal and inverse orientations), the four functional Jκ segments, the intronic κ enhancer (Eik), and the constant Cκ exon. (right) In-frame (F) and out-of-frame (VPTC and CPTC) Vκ-Jκ5 junctions are depicted, and the position of normal stop codons (stop) and PTCs are shown. Primers used are indicated (black arrows). (B) Representation of in-frame (F), VPTC, and CPTC mini-gene constructs. Out-of-frame junctions were created by inserting 5 nt (VPTC) or 10 nt (CPTC) at Vκ-Jκ5 junctions. (C) RT-PCR was performed using L1-135-for/Cκ-rev primers to identify full-length (κLC) and alternatively spliced (ΔV-κLC) mRNAs simultaneously. (D) Semi-quantitative RT-PCR analysis of S194 cell lines transfected with F, VPTC, and CPTC constructs. (E) A20 (mature B) and S194 (PC) cell lines were cotransfected with F and either VPTC or CPTC constructs (A20-F+VPTC, n = 9; A20-F+CPTC, n = 5; S194-F+VPTC, n = 6; S194-F+CPTC, n = 5). Relative ΔV-κLC mRNA levels were quantified using capillary electrophoresis by dividing areas of peaks corresponding to ΔV-κLC mRNAs to those of control functional (F) κLC mRNAs (ΔV-κLC/F ratios) as previously described (Chemin et al., 2010); part of these data [A20-F+VPTC, n = 4; S194-F+VPTC, n = 3] comes from this previous study. Data are representative of at least three independent experiments (n = 5–9). Bars represent mean expression ± SEM. Unpaired two-tailed Student’s test was performed to determine significance. ***, P < 0.001.
the selection of terminally differentiated cells, thus providing indirect arguments for a TIE checkpoint. To address this issue, we analyzed the Igκ repertoire at the DNA level in B cells and PCs from WT animals. We determined CDR3 lengths and distinguished in-frame (F or VJ+) VPTC and CPTC VκJκ junctions (Fig. 1 A). In agreement with previous works (Chen et al., 1993; Ehlich et al., 1993; Novobrantseva et al., 1999; Bertocci et al., 2003; Delpy et al., 2004a), VκJκ junctions were readily detectable in precursor B cells that did not express a BCR (pro–B/pre–B cells; B220b/Igλ−). As expected, numerous (∼66.6%) out-of-frame junctions were retrieved in the absence of BCR-mediated selection, with an equivalent repartition of VPTC and CPTC subclasses (Table 1). In contrast, the percentage of F junctions was much higher in GC B cells (B220b/PNAb) having been selected for functional BCR expression, with a reciprocal fall of VPTC and CPTC subclasses (Table 1). Unexpectedly, the frequency of VPTC junctions still more dramatically decreased in PCs (B220pos/CD138pos; approximately nine-fold lower than GC B cells; P < 0.01), whereas that of CPTC junctions was unchanged (Table 1). Next, we addressed whether the disappearance of PCs harboring VPTC-rearranged Igκ alleles prone to exon-skipping correlated with a decrease in ΔV-κLC mRNAs. ΔV-κLC mRNAs were analyzed by RT-PCR in sorted B cell populations and PCs (Fig. 3 A). Although, a short band corresponding to ΔV-κLC mRNAs was readily detectable in B cell populations, these alternative transcripts were not retrieved in PCs (Fig. 3, B and C). The complete loss of ΔV-κLC mRNAs in PCs was consistent with the drastic disappearance of VPTC junctions in their repertoire. Altogether, the selective elimination of PCs with biallelic Igκ rearrangements in VJ/VPTC configuration, likely reflects toxic effects of ΔV-κLCs. This suggests that, independently of B cell functionality and specificity for antigen, a late TIE checkpoint counterselects B lineage cells producing aberrant Ig and prevents their survival as terminally differentiated PCs.

**Impaired PC differentiation upon inducible expression of ΔV-κLCs**

According to previous observations showing that ∼40–50% of B lymphocytes exhibit biallelic V(D)J rearrangements of Ig genes (Mostoslavsky et al., 2004; Daly et al., 2007), approximately half of these B cell clones (∼20–25%), i.e., those harboring VPTC-rearranged Igκ alleles, are candidates for an Ag-independent TIE checkpoint, eventually leading to their elimination at the step of PC differentiation. However, the rarity of VPTC junctions in PCs at the DNA level is only an indirect proof that cells go through such a checkpoint. To directly address the impact of ΔV-κLCs on late B cell development and on humoral responses, we created Igκ knock-in mice allowing inducible expression of ΔV-κLCs in B lineage cells (Fig. 4 A). In these mutants, later referred as inducible TIE (iTIE) mice, the replacement of endogenous Jκ segments by a leader exon (L1-33) under the control of a pVH promoter led to enforced expression of ΔV-κLC mRNAs and proteins. Conditional Cre-mediated expression of ΔV-κLC mRNAs was achieved by inserting a loxP-flanked neomycin-resistance (NeoR) cassette that blocks transcription and splicing of the L1-33 exon (Fig. 4 A). In heterozygous (iTIE+/+) animals, the WT allele (+) supports normal Igκ expression during B

![Figure 2](image-url)
cell development, whereas the knock-in (iTIE) allele allows Cre-dependent expression of ΔV-κLCs. After mating iTIE/+ mice with a CMV-Cre–expressing strain, high ΔV-κLC mRNA levels and significant truncated Ig synthesis were observed in Cre-positive (Crepos) B lineage cells, with no major leakiness in Cre-negative (Creneg) controls (Fig. 4 B). In agreement with the aforementioned data from transfected cells (Fig. 2), ΔV-κLC amounts increased after treatment with the proteasome inhibitor bortezomib (Bz; Fig. 4 B). Interestingly, these knock-in mice mimic the outcome of the physiological TIE checkpoint, as the production of ΔV-κLCs severely impairs PC differentiation without perturbing B cell development and GC formation (Fig. 4, C and D). In controls, the sole expression of Cre-recombinase affected neither total B220pos and GC B cells nor PCs (unpublished data). Consistent with a global defect in PC differentiation, Crepos iTIE/+ mice exhibited significantly lower T cell–independent and T cell–dependent antibody responses than Crepos iTIE/+ controls (Fig. 4, E and F). These findings were further confirmed in competitive experiments mimicking the occurrence of the TIE checkpoint during late B cell differentiation and in only a fraction of terminally differentiated PCs. For that, we analyzed B cell and PC contents after mating iTIE mice with AID-Cre-EYFP mice, allowing conditional Cre-recombinase expression in GC B cells upon treatment with tamoxifen (Dogan et al., 2009). We found significantly lower amounts of

Table 1. Analysis of productive and nonproductive VκJκ5 junctions in B and plasma cells from WT mice

| Cell type       | Total | In-frame | VPTC | CPTC |
|-----------------|-------|----------|------|------|
| Pro–B/pre–B cells | 51    | 33.3     | 31.4 | 35.3 |
| GC B cells      | 53    | 60.4     | 18.9 | 20.7 |
| Plasma cells    | 47    | 72.3     | 2.1  | 25.5 |

Pro–B/pre–B cells (B220pos/IgLneg) and PCs (B220pos/CD138pos) were isolated from bone marrow and GC B cells (B220pos/PNAhi) from Peyer’s patches, and PCR were performed on genomic DNA using Vκ cons-for/Jκ5-rev primers (Fig. 1 A). Productive VκJκ5 junctions were assigned as in-frame and nonproductive junctions as VPTC and CPTC. Data are from two independent cell sorting experiments. n = total number of sequences.
EYFP-positive PCs in AID-Cre-EYFP iTIE/+ mice, compared with AID-Cre-EYFP +/- littermates (Fig. 5, A and B). In contrast, similar frequency and absolute numbers of EYFP-positive B cells were retrieved in these animals (Fig. 5, C and D). Thus, the PC defect observed in Cre<sup>enh</sup> iTIE mice confirms that the production of ΔV-κLCs induces deleterious effects in terminally differentiated B lineage cells.

**Short lifespan and low levels of Ig secretion in PCs expressing ΔV-κLCs**

Next, we sought to elucidate how these truncated Ig control plasma cell maturation using the iTIE mouse model with enforced ΔV-κLC expression. Experiments were performed in homozygous iTIE/iTIE animals that have no functional Igκ alleles and exhibit only Igλ-expressing B lineage cells. Like
heterozygous mutant mice (Cre<sup>pos</sup> iTIE/+), but to a higher extent, homozygous animals (Cre<sup>pos</sup> iTIE/iTIE) exhibited very low amounts of B220<sup>neg</sup>CD138<sup>pos</sup> PCs (Fig. 6 A) together with a strong reduction in serum Ig levels (Fig. 6 B), compared with Cre<sup>neg</sup> controls. ΔV-κLC mRNA amounts were determined by qPCR in B cells and PCs isolated from bone marrow of Crepos iTIE/iTIE mice. We found that the remaining PCs exhibited approximately five-fold higher ΔV-κLC mRNA levels compared with sorted B cells (Fig. 6 C). To evaluate whether ΔV-κLC production in PCs affected Ig synthesis, we analyzed the expression of Igκ in Cre<sup>pos</sup> iTIE/iTIE mice. Interestingly, we found an approximately two-fold lower mean fluorescence intensity (MFI) for intracellular Igκ in Cre<sup>pos</sup> iTIE/iTIE PCs, compared with Cre<sup>neg</sup> iTIE/iTIE counterparts (Fig. 6 D). Hence, PCs expressing ΔV-κLCs synthesized low levels of Ig. Next, we evaluated whether the truncated Ig impaired the renewal or survival of PCs in Cre<sup>pos</sup> and Cre<sup>pos</sup> iTIE/iTIE mice injected with BrdU. Similar amounts of BrdU<sup>pos</sup> and BrdU<sup>neg</sup> B cells were retrieved for both strains (Fig. 6 E). In contrast, the frequency of BrdU<sup>pos</sup> (cycling) B220<sup>pos</sup>CD138<sup>pos</sup> cells was significantly increased in Cre<sup>pos</sup> iTIE mice compared with Cre<sup>pos</sup> controls, indicating higher proportion of recently divided plasmablasts (PBs) upon production of ΔV-κLCs. Inversely, we found lower amounts of noncycling BrdU<sup>pos</sup> PCs in Cre<sup>pos</sup> iTIE mice. Next, we sought to elucidate how the production of truncated Ig limited the lifespan of PCs. We analyzed the expression of ER stress markers, together with unfolded protein response (UPR) components in sorted PCs (B220<sup>pos</sup>CD138<sup>pos</sup>). In agreement with an exacerbated ER stress response upon production of ΔV-κLCs, Chop, homocysteine-induced ER protein (Herp), X-box binding protein 1 spliced (Xbp1s), and BiP mRNA levels were strongly increased in PCs isolated from Cre<sup>pos</sup> iTIE mice (Fig. 6 F). Thus, the rise of ER stress as a result of the production of ΔV-κLCs is transiently tolerated in early differentiating PBs but clearly compromises their survival as noncycling long-lived PCs secreting high Ig amounts (Nutt et al., 2015).

Intrinsic toxicity of truncated Ig in plasma cells
ΔV-κLCs can induce multiple deleterious effects. On one hand, their presence could impede the normal assembly of IgH and IgL chains and relate to the classical quality control of BCR assembly (Melchers et al., 2000). On the other hand, the TIE checkpoint could involve an intrinsic toxicity of truncated Ig independent of BCR expression. To distinguish between these two hypotheses, we determined whether the elimination of PCs containing V<sub>PTC</sub>-rearranged Igκ alleles also occurred in the absence of IgH chain expression. Igκ repertoires were thus compared between B cells (B220<sup>pos</sup>) and PCs (B220<sup>neg</sup>/CD138<sup>pos</sup>) isolated from spleens of DH-LMP2A mice (Lechouane et al., 2013). In this model, the Epstein-Barr virus LMP2A protein mimics the BCR tonic signal and B lymphocytes develop without any IgH chain (Casola et al., 2004). As expected upon such a by-pass of BCR-driven positive selection, approximately two thirds of VκJκ5 junctions were out-of-frame in DH-LMP2A B cells, with an equivalent reparation of V<sub>PTC</sub> and C<sub>PTC</sub> subclasses (Table 2). Regarding nonproductive junctions, we found again that the V<sub>PTC</sub> class was drastically decreased in PCs, compared with B cells (Table 2). In addition, we confirmed that the sole presence of ΔV-κLCs impaired PC differentiation by breeding iTIE mice on a DH-LMP2A background (Fig. 7 A). Thus, the elimination of PCs containing V<sub>PTC</sub>-rearranged Igκ alleles occurred independent of IgH expression and BCR assembly. According to an exacerbated ER stress response and activation of the UPR, CHOP, HERP, BiP, Xbp1s, and IRE1α expression were strongly increased upon production of ΔV-κLCs (Fig. 7).
Table 2. Analysis of productive and nonproductive VκJκ5 junctions in B and plasma cells from DH-LMP2A mice

| Cell type         | Total | In-frame | VPTC % | CPTC % | CPTC % |
|-------------------|-------|----------|--------|--------|--------|
| B cells           | 53    | 35.8     | 32.1   | 32.1   |        |
| Plasma cells      | 55    | 67.3     | 27.3   | 5.4    | 27.3   |

B cells (B220<sup>+</sup>/CD138<sup>-</sup>) and PCs (B220<sup>-</sup>/CD138<sup>+</sup>) were isolated from spleens of DH-LMP2A mice and PCRs were performed on genomic DNA using Vκ cons-for/Jκ5rev primers (Fig. 1A). Productive and nonproductive VκJκ5 junctions were assigned as in-frame, VPTC, and CPTC junctions, as described in Table 1. Data are from two independent cell sorting experiments. n = total number of sequences.
Likewise, high ΔV-κLC mRNA levels were correlated with a strong expression of ER stress markers, in PBs sorted 4 d after LPS stimulation (Fig. 7, D and E). Altogether, these findings indicate that the TIE checkpoint is unrelated to a disturbed IgH/IgL assembly, but rather to a toxic ER stress provoked by the sole presence of ΔV-κLCs.

DISCUSSION

The allelic exclusion of Ig genes is stringently established before or during early B cell maturation and ensures the monospecificity of B lymphocytes through stepwise V(D)J recombination of Ig alleles. In cells harboring biallelic V(D)J rearrangements, the noise coming from transcription of nonproductive alleles has long been neglected because these transcripts are either degraded by NMD or do not encode functional Ig chains (Jäck et al., 1989; Li and Wilkinson, 1998; Delpy et al., 2004a; Chemin et al., 2010; Tinguely et al., 2012). However, we report here that alternative splicing of these passenger transcripts leads to the production of aberrant Ig chains that seriously impede PC differentiation. These findings identify a novel TIE checkpoint that eliminates ∼20–25% of terminally differentiated B lineage cells, i.e., those transcribing nonproductive Igκ alleles prone to exon skipping. Consistent with an Ag-independent checkpoint, ΔV-κLCs exert intrinsic toxic effects in PCs, likely mediated by ER stress–induced apoptosis. In addition, PCs expressing ΔV-κLCs synthesize low amounts of Ig and are mostly found among short-lived PBs. We propose that the TIE-checkpoint favors the selection of long-lived PCs with limited basal ER stress supporting high levels of Ig secretion.

Regarding nonproductively rearranged Igκ alleles, VPTC and CPTC junctions are highly similar and often exhibit a single nucleotide difference within the CDR3 sequences. Despite minor changes at the DNA level, we found that exon skipping was only enhanced for VPTC class transcripts, with higher magnitude in PCs than in mature B cells. Although the molecular mechanisms underlying NAS are not completely understood, this response is triggered by two distinct pathways referred as class I and class II NAS (Maquat, 2002; Wang et al., 2002; Chang et al., 2007). Class I NAS occurs upon disruption of splicing motifs, such as exonic/intronic splicing enhancers/silencers, by either nonsense or missense mutations, whereas class II NAS is a reading frame–dependent process strictly induced by nonsense mutations (Chang et al., 2007). Thus, our data strongly suggest that a class II NAS response occurs for nonproductive Igκ transcripts that
further relies on PTC recognition within the skipped V exon, but not within the downstream Cκ exon. We also observed an approximately two-fold increase in ΔV-κLC mRNA levels in S194 plasma cells, compared with A20 B cells. In PCs, Ig genes are localized in transcription factories near the nuclear pore, and these specialized areas authorize cooperation of enhancers and facilitate Ig gene transcription (Park et al., 2014). In addition, a rise of exon skipping has been observed for highly transcribed genes, leading to the assumption that alternative splicing is closely correlated to the rate of RNA polymerase II elongation (Nogués et al., 2003; Shukla and Oberdoerffer, 2012). Altogether, we suggest that the boost of Ig gene transcription accompanying PC differentiation detrimentally promotes exon skipping during splicing of non-productive Igκ premRNAs.

Biallelic Igκ rearrangements with a VJ+/VPTC configuration represent around 20–25% of mature B cells (Mostoslavsky et al., 2004) and hereby yield the population affected by the TIE checkpoint in terminally differentiated PCs. Sensitivity to prolonged ER stress likely explains the elimination of PCs producing ΔV-κLCs. Accordingly, ER stress markers and UPR components such as CHOP, BIP, HERP, and XBP1/IRE1α are up-regulated upon production of ΔV-κLCs in PCs. In addition, we found that Chop mRNA levels were already elevated in B cells expressing truncated Ig, suggesting that the early expression of this pro-apoptotic factor could influence the fate of differentiating PCs. Although a previous study showed that CHOP is dispensable for the development of LPS-induced plasmablasts (Masciarelli et al., 2010), our data are in agreement with a recent work highlighting its role in differentiating PCs (Gaudette et al., 2014) and also demonstrating that BCL-XL expression protects PCs from CHOP-associated apoptosis. The iTIE mouse model described herein should be useful to decipher the complex interplay between pro- and anti-apoptotic factors during PC differentiation.

With regard to PC dyscrasias, truncated Ig have been observed in some cases of myeloma (Cogné et al., 1988, 1992; Cogné and Guglielmi, 1993). However, it is a rare feature of primary PC dyscrasias and of malignant PC lines, suggesting that the TIE checkpoint also shapes the human Ig repertoire and that PCs producing aberrant Ig molecules are rarely rescued by oncogenic events (Decourt et al., 2004). Accordingly, we found that ΔV-κLCs were harmful for survival of normal PCs but exerted no obvious effect when expressed in tumor PCs in the absence of PI treatment. The survival of normal and malignant PCs is influenced by ER stress and the balance between load versus capacity of the proteasome (Cenci and Sitia, 2007). Being able to modulate ER stress and proteasome activity is of considerable interest for multiple myeloma (MM) patients (Meister et al., 2007) and new therapeutic strategies have emerged combining inhibitors of chaperones (Hsp90: Heat shock protein 90) or factors involved in unfolded protein response (IRE1α) with PI (Ishii et al., 2012; Mimura et al., 2012). Our findings are consistent with these combined approaches, increasing the amount of aberrant proteins while decreasing their proteasomal degradation, and suggest that splicing modulators or NMD inhibitors could also be useful tools to reinforce the production of truncated proteins including ΔV-κLCs.

Interestingly, the late TIE checkpoint stands as a quality control of individual and unassembled ΔV-κLCs and is not a result of a disturbance of BCR or secreted Ig assembly (which could then be viewed as a failed Ag response). Thus, the late occurrence of the TIE checkpoint, at the terminal differentiation stage, can be considered an intrinsic waste and weakness of the immune system. It allows uncensored B cells to accumulate as mature B cells carrying VPTC-rearranged Igκ alleles, to potentially respond to Ag, and undergo affinity maturation in GCs, while being unable to yield long-lived PCs secreting to potentially respond to Ag, and undergo affinity maturation in GCs, while being unable to yield long-lived PCs secreting high amounts of Ig (Nutt et al., 2015). The TIE checkpoint then cuts off the diversity of PCs arising from Ag-stimulated B cells for reasons unrelated to Ag recognition and affinity.

Whereas NMD basically protects cells from truncated protein synthesis, we show that activation of NAS herein exerts an opposite effect, finally ending with loss of ΔV-κLC-expressing PCs. This study thus shows that transcription and splicing of non-productive Ig alleles, although often neglected or considered to be under control, are in fact often inappropriately handled by RNA surveillance pathways, allows production of truncated Ig and ends with a PC wastage.

**MATERIALS AND METHODS**

**Gene targeting.** Homologous recombination at the Igκ light chain locus was performed as described previously (Sirac et al., 2006). In brief, a 12.7-kb BamHI genomic fragment corresponding to the germline mouse JκCκ cluster was used to generate the iTIE targeting construct (Van Ness et al., 1982). A 2.2-kb BsmI–SacII fragment spanning all the Jκ segments was replaced with a cassette containing a Vκ promoter (pVκ), a human leader exon (L1,33), and a hsvTk-flanked neomycin-resistance gene driven by the Herpes simplex thymidine kinase promoter (hsvTk-neoR). The hsvTk-neoR cassette was inserted in opposite orientation to block the transcription and/or splicing of the L1,33 exon (Fig. 4A). Mouse embryonic stem cells (E14) were transfected with the linearized vector by electroporation and selected using 300 μg/ml Geneticin and 2 μM ganciclovir.

**Mice.** 2–3-mo-old mice were used in all experiments and maintained in our animal facilities, at 21–23°C with a 12-h light/dark cycle. Experiments were performed according to the guidelines of the ethics committee in Animal Experimentation of Limousin (registered by the National Committee under the number C2EA-33) and were approved as part of the protocol registered under the number CREEL 6–07–2012. Heterozygous mutant mice (iTIE+/+) were backcrossed to C57BL/6 (B6) for at least three generations, and then mated with Cre-expressing mice to induce deletion of the hsvTk-neoR cassette. B6 CMV-Cre mice were obtained.
from the Mouse Clinical Institute (Illkirch, France). B6 AID-Cre-EYFP mice, harboring a tamoxifen-inducible Cre recombinase enzyme controlled by the Aicda (activation-induced cytidine deaminase) promoter and a loxP-flanked EYFP (enhanced yellow fluorescent protein) reporter gene, have been described elsewhere (Dogan et al., 2009). B6 AID-Cre-EYFP mice were obtained from the Imagine Institute (Paris, France). In DH-LMP2A mice (with mixed B6/BALB/c backgrounds), the Epstein-Barr virus LMP2A protein drives B cell development and plasma cell differentiation, as previously described (Casola et al., 2004; Lechouane et al., 2013). DH-LMP2A mice were obtained from the Institute of Molecular Oncology Foundation (Milano, Italy).

**Cell transfection.** A20, S194, and Sp2/0 murine cell lines were cultured (10^6 cells/ml) in RPMI supplemented with 10% fetal calf serum (Invitrogen), sodium pyruvate, nonessential amino acids, β-mercaptoethanol, 100 U/ml penicillin, and 100 μg/ml streptomycin (Gibco). Cells (2 × 10^6) were stably transfected by electroporation according to the manufacturer’s instructions (Amaxa). Transfections were performed using in-frame (F), V^PTC^ and C^PTC^ linear plasmid constructs separately, or using equimolar ratios of F and nonproductive constructs as previously described (Chemin et al., 2010). V_kappajunctions were built by artificially joining the V_kappa,135 and J_kappa segments. To create a reading frameshift, 5 and 10 additional nt were introduced at the V_kappa junction in V^PTC^ and C^PTC^ constructs, respectively (Fig. 1 B). Cells (10^6 cells/ml) were treated or not with MG132 (1 µg/ml) for 5 or 8 h, as indicated, and control cells were treated with DMSO alone (dilution factor 1/1,000).

**Flow cytometry and cell sorting.** The frequency of apoptotic cells was determined after staining with Annexin V (BD) and propidium iodide using a LSRII Fortessa (BD). Data were analyzed with FACSDiva software (BD). Bone marrow B cell precursors and PCs were sorted on a FACSVantage (BD) after staining with anti-mouse B220 (RA3-6B2; BioLegend), anti–mouse Igκ (187–1; Beckman Coulter), anti–mouse Igκ (JC5–1; Beckman Coulter), and anti–mouse CD138 (281–2; BD) mAbs. The gates used for cell sorting are indicated in Fig. 3 A. GC B cells were isolated from Peyer’s patches after staining with anti–mouse B220 and PNA (peanut agglutinin; Sigma-Aldrich) as previously described (Delpy et al., 2004b). In all experiments, purity of sorted cells was >90%. B cells were also isolated from spleens by negative selection using anti–CD43 microbeads (Miltenyi Biotec) and stimulated (0.5 × 10^6 cells/ml) with 1 µg/ml of LPS (LPS-EB UltraPure; InvivoGen) for 4 d.

To determine the frequency of GC B cells and PCs, erythrocyte-depleted spleen cells were stained with anti–mouse GL7 (GL7; BD), anti-B220, and anti–CD138 mAbs, 7 d after intraperitoneal (IP) injection of sheep red blood cells (SRBCs; bioMérieux). When indicated, mice received additional subcutaneous bortezomib (Sillag) injections at day 5 and 6 (0.5 mg/kg). To induce nuclear translocation of Cre-recombinase, AID-Cre-EYFP mice were treated with Tamoxifen (1 µg/mouse; IP; Sigma-Aldrich) at day 2, 4, and 6 after SRBC immunization.

To analyze the renewal of B lineage cells in bone marrow, mice received i.p. BrdU (Sigma-Aldrich) injections (1 mg/mouse at day 0, then 0.5 mg every 48 h) for 10 d. Cells were then stained according to the BrdU Flow kit (BD) protocol. Cells were treated with DNase I before staining with anti-BrdU mAb. Intracellular Igκ staining was performed in bone marrow cells isolated from homozygous iTIE/iTIE mice using the Cytofix/Cytoperm kit (BD).

**Western blot.** For Western blot analysis, a 4–20% Mini-PRO TEAN TGX polyacrylamide gel (Bio-Rad Laboratories) was used. Each sample was then denatured at 94°C for 3 min before being loaded. Gels were blotted onto TransBlot Turbo polyvinylidene fluoride membranes (Bio-Rad Laboratories), and blocked in PBST buffer (137 mM NaCl, 2.7 mM KCl, 10 mM Na₂HPO₄, 2 mM KH₂PO₄, and 0.1% Tween 20, pH 7.4) containing 5% nonfat dry milk. The signal was measured by chemoluminescence (ECL plus; GE Healthcare). Western blots were performed using rabbit anti-mouse HERP (Santa Cruz Biotechnology, Inc.), CHOP, BIP, IRE1α (Cell Signaling Technology), goat anti–mouse Igκ (Beckman Coulter), and using goat anti–mouse GAPDH (R&D Systems) Abs for normalization.

**ELISA assays.** Ig titers were determined in culture supernatants and mouse sera using unlabeled and alkaline phosphatase–conjugated goat anti–mouse Abs (IgM, IgG1, IgG3, IgG2b, Igκ, Igλ, and total IgG; Southern Biotech) as described (Pinaud et al., 2001; Sirac et al., 2006). Total Ig amounts were revealed by the addition of p-nitrophenyl phosphate (Sigma-Aldrich), and plates were read at 405 nm.

To analyze T cell–dependent humoral responses, mice received two IP injections with 10 μg hen egg OVA (Sigma-Aldrich) emulsified in 50% CFA (day 0) and IFA (day 14). For T cell–independent responses, mice received two IP injections (days 0 and 14) with 50 μg of 2,4, Dinitrophenyl-Amino-Ethyl-Carboxy-Methyl-Ficoll (DNPE-Ficoll; Biosearch Technologies) emulsified in 50% IFA. Blood samples were collected (day 7 and 24) and antigen–specific antibody titers were determined in polycarbonate 96 multiwell plates (Maxisorp; Nunc) coated overnight at 4°C with OVA or albumin-DNP (Sigma-Aldrich) solution (10 μg/ml), in carbonate buffer.

**PCR and RT-PCR.** Genomic DNA and total RNA were prepared using standard protease K (Eurogentec) and Tri-reagent (Invitrogen) procedures, respectively. RT-PCR was performed on DNase I-treated (Invitrogen) RNA and was negative in the absence of reverse transcription, ruling out contamination by genomic DNA. Reverse transcription was performed using Superscript II (Invitrogen) on 1 to 5...
VPTC, and CPTC Vκ junctions were analyzed after cloning of κ sequences of VJκ. Identification of PCR products was also performed using an Invitrogen, using V-QUEST software (IMGT, the international ImMunoGeneTics information system). Real-time PCR were performed on a ABI PRISM 7000 Sequence Detection System (Life Technologies). Transcripts were quantified according to the standard 2−ΔΔCt method after normalization to Gapdh. Sequences of primers and probes are available upon request.

**Statistical analysis.** Results are expressed as mean ± SEM and overall differences between variables were evaluated by a two-tailed unpaired Student’s t test using Prism GraphPad software. A χ² test was done to analyze the distribution of F, VPTCκ, and CPTC Vκ junctions.

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