Endogenous Retroviruses Transcriptional Modulation After Severe Infection, Trauma and Burn

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Although human endogenous retroviruses (HERVs) expression is a growing subject of interest, no study focused before on specific endogenous retroviruses loci activation in severely injured patients. Yet, HERV reactivation is observed in immunity compromised settings like some cancers and auto-immune diseases. Our objective was to assess the transcriptional modulation of HERVs in burn, trauma and septic shock patients. We analyzed HERV transcriptome with microarray data from whole blood samples of a burn cohort (n = 30), a trauma cohort (n = 105) and 2 septic shock cohorts (n = 28, n = 51), and healthy volunteers (HV, n = 60). We described expression of the 337 probesets targeting HERV from U133 plus 2.0 microarray in each dataset and then we compared HERVs transcriptional modulation of patients compared to healthy volunteers. Although all 4 cohorts contained critically ill patients, the majority of the 337 HERVs was not expressed (around 74% in mean). Each cohort had differentially expressed probesets in patients compared to HV (from 19 to 46). Strikingly, 5 HERVs were in common in all types of severely injured patients, with 4 being up-modulated in patients. We highlighted co-expressed profiles between HERV and nearby CD55 and CD300LF genes as well as autonomous HERV expression. We suggest an inflammatory-specific HERV transcriptional response, and importantly, we introduce that the HERVs close to immunity-related genes might have a role on its expression.

Keywords: endogenous retroviruses, severe inflammatory injuries, septic shock, burn, trauma, transcriptome, host response
INTRODUCTION

Human Endogenous Retroviruses (HERVs) are former exogenous retroviruses which have infected germinal cells and became integrated in our genome million years ago (1). These rare events happened several times in evolution. As retrotansposons, they are able to duplicate across the genome and they represent today more than 8% of our genome. Each insertion therefore led to distinct groups or families, each including multiple copies. Current classification annotates around 100 such groups.

HERV loci initially shared a common structure with exogenous retroviruses: internal protein coding regions (gag, pro, pol, env) flanked by two identical Long Terminal Repeats (LTRs). The accumulation of mutations and recombination events during evolution made most of these elements incomplete and defective for replication. Indeed, most of HERVs in our genome are now solo LTRs (1) resulting from recombination between 5′ and 3′ proviral LTRs. LTRs are critical elements that control viral gene expression either as promoters, enhancers or as polyadenylation signals. When inserted upstream, within or downstream of a “conventional” protein coding gene, LTRs can modulate its expression pattern (2, 3). For example, the presence of intronic LTR can result in novel transcripts, by providing alternative promoters, enhancers or polyadenylation signals, or by altering RNA splicing (4–6). Very few is known about of the transcriptional modulation of such elements in pathological contexts but in cancers [like testicular cancer (7) or colorectal cancer (8)] and auto-immune diseases [like multiple sclerosis (9–11)].

Few studies focused on HERVs reactivation in acute inflammatory contexts. In mice, modulation of HERV’s expression has been shown to be quite specific, with signatures related to pathogen-associated molecular pattern (PAMPs) (12). In human, LPS or PMA stimulations of myeloid cells revealed an increase expression of four HERVs families (13). In vivo, HERVs expression has been detected in the plasma and whole blood samples of burn patients (14, 15) although the studies focused on whole HERVs families, not on specific loci. Studying HERV transcriptome modulation after severe inflammatory injuries could help to better understand pathological states of patients.

After severe injuries like septic shock, burn or trauma, leading to an important inflammatory response, we and others have shown that the blood transcriptome is highly modulated, with early and profound changes in adaptive and innate immune responses (16, 17). Moreover, in these contexts, viral reactivation is often observed, especially for Herpes Viruses (18, 19). This reactivation is associated with an immunosuppressive state (20). We therefore hypothesize that HERV, like latent viruses, may reactivate and be transcribed in vivo after inflammatory injuries. Given that several groups showed that some probes of commercial whole genome microarray do target HERV loci (21, 22) (such as Affymetrix U133 plus 2), we retrospectively explored microarray datasets obtained in our lab to study the HERV transcriptome modulation in various contexts of injuries in vivo.

MATERIALS AND METHODS

Patients and Sample Collection

Microarray Analyzed Cohort

BURNS COHORT

30 severe burn patients admitted at Hospices Civils de Lyon, France (HCL) were included in a placebo-controlled, randomized, double-blind study assessing the efficacy of hydrocortisone administration on burn shock duration. Inclusion / exclusion criteria, clinical description and ethical considerations of the cohort have been previously published elsewhere (16, 23). Thirteen healthy volunteers were also recruited within Hospices Civils de Lyon to serve as controls for the transcriptional study. Whole blood samples were collected at inclusion (severe shock, before any treatment, Day 1) and in the following days (around day 2 (D2), day 5 (D5), and day 7 (D7) after inclusion).

TRAUMAS COHORT

105 patients with severe trauma were admitted at HCL. Briefly, patients were included when they were under mechanic ventilation, with an Injury Severity Score (ISS) over 25 and were at least 18 years old. Inclusion / exclusion criteria and ethical considerations of the cohort have been previously published elsewhere (24). The main clinical variables are summarized on Table S1. Samples were collected at day 1 (D1) or day 2 (D2) after trauma. Data from 22 healthy volunteers were also used to make comparisons with patients (identical with septic shock cohort 2).

Septic shock cohort 1 (SS1)

Twenty-eight septic shock patients and 25 HV admitted into 2 ICUs of HCL were included in this study to explore the early transcriptome modulation after septic shock. Inclusion/exclusion criteria, clinical description and ethical considerations of the cohort have been previously published elsewhere (25). The first blood sample was collected at the onset of shock (i.e., within 30 min after the beginning of vasoactive treatment, D0) and at day 1 (D1) and day 2 (D2) after shock.

Septic shock cohort 2 (SS2)

Fifty-one septic shock patients and 25 HV admitted into 2 ICUs of HCL and 22 HV were included in a prognostic biomarker study. Inclusion / exclusion criteria, clinical description and ethical considerations of the cohort have been previously published elsewhere (26). Samples were collected at day 1 (D1), day 2 (D2) and day 3 (D3) after shock.

RT-qPCR Validation Cohorts

Patients

Subset of cohorts used for microarray analysis were used for validation cohort: 10 burn samples at D1, 10 traumas samples at D1, 10 SS1 samples at D1, 10 SS2 samples at D1. Each subset was
matched with its corresponding cohort: Age, sex and Total Burn Surface Area (TBSA) for burns-Sex, Sepsis at D7 and Death at D28 for traumas-Age, sex and SAPS II for SS1-Age, Sex and Death at D28 for SS2.

**Healthy Volunteers**
Whole blood samples were purchased from the Etablissement Français du Sang \( (n = 12) \). The mean age of HV is 56, with a standard error of 9. According to the standardized procedure for blood donation, written informed consent was obtained from healthy volunteers (HVs) and personal data for blood donors were anonymized at time of blood donation and before blood transfer to a research lab.

**Flow Cytometry Validation Cohort**

**Burns**
Whole blood samples (EDTA tubes) from 13 burn patients sampled at D1 and D7 and admitted in Edouard Herriot hospital at Lyon, France were recruited as part of the EARLYBURN study (NCT02940171). Patients were aged from 21 to 84 (mean = 53), 12 men. The mean TBSA was 33\% (from 20 to 52\%). All samples from these patients were used for CD300LF protein analysis, and 7 of these 13 patients were used for CD55 protein analysis.

**Septic shocks**
Whole blood samples (EDTA tubes) from 22 septic shock patients sampled at D1/D2, D3/D4/D5, and D6/D7/D8 after shock and admitted in Edouard Herriot hospital at Lyon, France were recruited as part of IMMUNOSEPSIS study (NCT02803346). Patients were aged from 23 to 81 (mean = 68), 16 men. Eleven samples were used for CD300LF protein analysis and 11 other samples for CD55 protein analysis.

**Healthy volunteers**
Whole blood samples (EDTA tubes) were purchased from the Etablissement Français du Sang \( (n = 18) \). Donors were aged from 21 to 63 (mean = 50), 12 men and 6 women. They were age-matched with burn and septic shock cohorts. According to the standardized procedure for blood donation, written informed consent was obtained from healthy volunteers (HVs) and personal data for blood donors were anonymized at time of blood donation and before blood transfer to a research lab.

**RNA Extraction and Microarrays**

Total RNA was extracted with PAXgene™ Blood RNA kit (PreAnalytiX, Hilden, Germany). Whole blood from PAXGene™ tubes was preferred to either buffy coat or PBMCs to ensure reproducibility and avoid missing samples within the context of a clinical study. RNA integrity was assessed using Agilent 2100 Bioanalyzer (Agilent Technologies, Waldbrum, Germany) and Lab-on-chip RNA 6000 Nano Assay (Agilent Technologies). Double-stranded cDNA was prepared from total RNA and an oligo-dT primer using GeneChip One-Cycle cDNA Synthesis Kit (Affymetrix, Santa Clara, United States). Three microgram labeled cRNA were hybridized onto Human Genome U133 Plus 2.0 GeneChips (Affymetrix), revealed and washed using FS450 fluidic station. GeneChips were scanned using a 5G scanner (Affymetrix) and images (DAT files) were converted to CEL files using GCOS software (Affymetrix).

**Microarray Analysis**

Microarray data are available on the Gene Expression Omnibus (GEO) website for Burn [GEO:GSE77791], SS1 [GEO:GSE57065], and SS2 [GEO:GSE95233] cohorts. The preprocessing methods were comparable in all datasets. Microarray normalization and statistical analysis were performed using R/Bioconductor (R v3.2.3). Quality assessment was performed through simpleaffy (v2.46.0) (27). After removing outlier samples the raw data were normalized, adjusted for background noise and summarized using the GCRMA (Guanine Cytosine Robust Multi-Array) algorithm with default parameters (28). COMBAT algorithm (29) was used to remove batch effect on Burn and Trauma cohorts. The 337 probesets from the U133 Plus2.0 microarray targeting HERVs have been identified and selected as described elsewhere (21, 22).

All the analysis were made with R (3.2.3). The differential expression analysis was performed with Limma package (3.26.9) (30). A probeset was considered significantly statistically differentially expressed between two conditions when absolute log2 Fold Change was higher than 0.5 and adjusted P-values [Benjamini-Hochberg correction (31)] lower than 0.01.

**Reverse Transcription and Quantitative PCR**

RNA from the cohorts, according to the above criteria, and new RNA from HV were selected. RNA concentration was determined using Quant-iT RNA, BR assay on Qubit (Life Technologies, Chicago, Illinois, United States). RNA integrity was assessed with the RNA 6000 Nano Kit on a Bioanalyzer (Agilent Technologies, Santa Clara, California, United States). Samples with RNA integrity number ≤6 were excluded due to poor quality RNA. Total RNA was reverse transcribed in complementary DNA (200 ng in a final volume of 20 µL) using QuantiTect Reverse Transcription kit (Qiagen) as recommended by the manufacturer. The expression levels of genes (CD55, CD300LF, SLC8A1, NFE4, PTTG1I2, and HPRT1 as reference gene) and associated HERVs were quantified using quantitative-real time polymerase chain reaction (qPCR). qPCR were performed on a LightCycler instrument using Light Cycler 480 Probes Master for the genes and reference genes and on SYBR Green I master for HERVs. Final volume of 20 µL contains 0.5 µM of primers. For genes, an initial denaturation step of 10 min at 95°C followed by 45 cycles, 10 s at 95°C, 29 s annealing at 60°C, and 1 s extension at 72°C, Taqman) was performed. For HERVs, an initial denaturation step of 5 min at 95°C followed by 45 cycles of a PCR protocol (10 s at 95°C, 15 s at 55°C and 15 s at 72°C, SYBR Green program), melting curve protocol was performed. The Second Derivative Maximum Method was used with the LightCycler software (Release 1.5.1) to automatically determine the crossing point for individual samples. Standard curves were generated by using serial dilutions of cDNA standards prepared from purified PCR amplicons obtained with the corresponding primers (Table S2). Relative standard curves describing the PCR efficiency of selected targets were created and used to perform
efficiency-corrected quantification with the LightCycler Relative Quantification Software. Targets expression normalization was performed using a selected housekeeping gene (hypoxanthine phosphoribosyltransferase 1 [HPRT1, (32)]), and results were expressed as normalized concentration ratio.

### Flow Cytometry

**Sampling and Staining**

The following antibodies were used: anti CD14-BV510, anti CD3–BV421 and anti CD56–PECy7 from BD Biosciences; anti CD300LF-PE from BD Biosciences or anti CD55-APC from Biolegend; anti CD16-APC from BD Biosciences or anti CD16-FITC from Beckman Coulter (Miami, FL) and PE Mouse IgG1, κ Isotype Control from BD Biosciences or APC Mouse IgG1, κ Isotype Control from R&D System. Red blood cell lysis was performed using Versalyse lysing solution (Beckman Coulter). CD300LF and CD55 expression were measured using Navios flow cytometer (Beckman-Coulter). Results were analyzed with Kaluza software (Beckman-Coulter) as Medians of Fluorescence Intensity (MFI).

### Statistics

Wilcoxon signed rank tests were done for RT-qPCR and flow cytometry results, by comparison between HV and each cohort of patients, for each target.

### ETHICS APPROVAL AND CONSENT TO PARTICIPATE

EDTA blood tubes were obtained from EFS (Etablissement Français du Sang) and used immediately. In accordance with EFS standardized procedures for blood donation, written no-objection was obtained from healthy volunteers to use the blood for the research and personal data for blood donors were anonymized before blood transfer to our research lab.

Protocols of the discovery and validation cohorts were approved by local ethics committees. Non-opposition to inclusion in the protocols was systematically recorded from patients or next of kin.

### RESULTS

We studied the in vivo modulation of the HERV transcriptome in three clinical relevant models of acute inflammatory injury: a burn, a trauma and 2 septic shock cohorts. We analyzed expression from each cohort independently comparing patients with healthy volunteers. All cohorts included severely injured patients (Table 1). The 30 burn patients had a median total burn surface area (TBSA) of 70% and high severity scores [median Baux: 110, median Abbreviated Burn Severity Index (ABSI): 11]. The 105 trauma patients had a median Injury Severity Score (ISS) score of 34 and a median Simplified Acute Physiology Score II (SAPSII) of 44. The 28 septic shocks from SS1 cohort had a median SAPSII of 45 and a median Charlson score of 2. The 51 patients from SS2 cohort had a median SAPSII of 51.

As previously published (21, 22), we extracted data from 337 probesets targeting HERVs loci from the whole genome U133 plus 2.0 microarray datasets. Among them, a majority had low expression levels, within background levels (Supplemental Figure 1). Based on hierarchical clustering analysis, 64 probesets (19%) were expressed (i.e., above background) for burns, 60 probesets (18%) for traumas, 164 for septic shock 1 (49%) and 63 for septic shocks 2 (19%). The 25% most variant probesets (n = 84) across samples in each dataset revealed that several probesets were even highly expressed (Figure 1). In each dataset, the hierarchical clustering highlighted a clear difference between patients and HV, suggesting a modulation of HERV expression following injury. Interestingly, over these top 25% most variant probesets selected in each dataset (resulting of 127 distinct probesets), 44 (35%) were similarly modulated in the four datasets, and 102 (80%) in at least 2 datasets (Supplemental Figure 2).

| Variable         | Burn (n = 30) | Trauma (n = 105) | Septic shock 1 (n = 28) | Septic shock 2 (n = 51) |
|------------------|--------------|-----------------|-------------------------|-------------------------|
| Age, years       | 48 (39–55)   | 38 (25–54)      | 62 (54–76)              | 65 (53–74)              |
| Gender, women, n (%) | 8 (27%)     | 34 (32%)        | 9 (32%)                 | 18 (35%)                |
| Weight, kg       | 94 (77–104)  | 78 (67–92)      | –                       | –                       |
| TBSA (%)         | 70 (48–84)   | –               | –                       | –                       |
| Baux score       | 110          | –               | –                       | –                       |
| ABSI score       | 11 (10–12)   | –               | –                       | –                       |
| ISS score        | –            | 34 (29–41)      | –                       | –                       |
| SOFA score       | –            | 5 (1–7)         | 10 (9–13)               | 10 (8–12)               |
| SAPSII score     | –            | 44 (29–56)      | 45 (34–56)              | 51 (43–62)              |
| Secondary septic shock | 12 (40%)    | 29 (28%)        | –                       | –                       |
| ICU length of stay, days | 66 (22–89)   | 9 (5–17)        | 10 (5–14)               | –                       |
| D28 mortality, n (%) | 8 (27%)     | 4 (4%)          | 5 (18%)                 | 17 (33%)                |

TBSA, Total Burn Surface Area (severe patient > 30%); Baux score, Predictor of mortality due to severe burns (severe patient > 100); ABSI, Abbreviated Burn Severity Index (severe patient > 9); ISS, Injury Severity Score (severe patient > 15); SOFA, Sequential organ failure assessment score (severe patient ≥ 3); SAPSII, Simplified Acute Physiology Score II, (severe patient > 30). In each cohort, cumulative mortality has been assessed up to the 28th day after ICU admission.
ICI admission (Supplemental Figure 4 and Table S3). Taking into account the global profile for each probeset, 16 (28.6%) had higher expression in patients compared to HV and 40 (71.4%) were down-modulated in patients. Interestingly, 5 probesets were differentially expressed in all 4 datasets and 16 in at least 3 of them (Figure 2A). All 5 commonly modulated probesets had consistent expression profile across the 4 datasets. Four were over-expressed in patients compared to healthy volunteers (Figure 2B). The 5th probeset, down-modulated in all datasets, maps at multiple locations in the genome and was not considered in further analyses. Among the 4 remaining modulated probesets, 1 HERV from ERV24B_Prim-int family (236982_at), is within 2kb from the PTTG1IP gene and 3 are within a gene. A HERV from LTR33 family (230354_at) is within an intron of SLC8A1 gene. A HERV from MLT1H family (1556107_at) and one from LTR16B2 family (1559777_at) are located in the 3'UTR of CD55 and MIR3945HG genes respectively (Table 2).

Moreover, we selected 2 other probesets of interest (1553043_a_at and 1560527_at, Figure 2C). The first one targets a MLT1D HERV located in the 3'UTR of CD300LF. It was up-modulated in burn and SS2 cohorts. It had a strong up-modulation at D1 in burn patients compared to HV, decreasing over the first week toward HV expression level at D7 (Supplemental Figure 5). The second one targets a LTR101_Mam HERV located in a 3'UTR of a processed transcripts of NFE4 gene. It was differentially expressed in the 2 septic shock cohorts. This probeset had the highest log2FC among the 5 septic shock-specific modulated probesets.

To validate these transcriptional HERV modulations, we designed primers on the 6 described HERV loci above, and on nearby genes by RT-qPCR (Table S2). For each targeted region, we made multiple RT-qPCR designs. We identified several distinct patterns of expression comparing HERVs and nearby genes (Tables 2): (i) for PTTG1IP and MIR3945HG regions, we observed no or low signal from the HERV loci (data not shown), (ii) for SLC8A1 (Figure 3) and NFE4 (Figure 4) regions, we observed a high signal from HERVs elements, but no or lower signal on the genes, (iii) for CD55 (Figure 5) and CD300LF (Figure 6) regions, we observed a middle or high signal from both HERV loci and genes.

To better interpret the results, we extracted from Ensembl the genome annotation and showed in genomic context, the microarray and the RT-qPCR results of SLC8A1 (Figure 3), NFE4 (Figure 4), CD55 (Figure 5) and CD300LF (Figure 6) regions.
SLC8A1 has 11 known transcripts. All but one are located in 3′ of the LTR33 HERV element targeted by the 230354_at probe, which is located in the first intron of SLC8A1-204 transcript (Figure 3A). The up-modulation of the LTR33 element in septic shock patients observed on microarray was confirmed by RT-qPCR (Figures 2B, 3B). The up-modulation observed for other cohorts was not confirmed by RT-qPCR. The gene SLC8A1 was not expressed in patients or HV, as seen on various microarray probesets and confirmed by RT-qPCR (SLC8A1_gene, var210, var211-212).

NFE4 gene has 2 transcripts (Figure 4A) and only one is coding for a protein (NFE4-202). The LTR101 Mam HERV element, targeted by the 2560527_at probe, is located in 3′ UTR of NFE4-201, the non-protein-coding transcript. Although the same trends are observed between microarray and RT-qPCR, the up-modulation of the LTR101 Mam element observed in septic shock patients with microarray was not statistically significant in RT-qPCR (Figures 2C, 4B). There was low or no signal on designs targeting gene transcripts (NFE4_gene and NFE4_gene_var201).

CD55 gene has 11 transcripts. The MLT1H HERV element, targeted by the 1556107_at probe, is located in the 3′ UTR of CD55-211 transcript (Figure 5A). The HERV element overlaps the 3′ UTR of transcript CD55-211 and a long intergenic noncoding RNA (lincRNA, a class of long transcribed RNA molecules longer than 200 nucleotides and not coding for proteins) (Figure 5B). The up-modulation of MLT1H seen with microarray in the 4 cohorts was partially confirmed by RT-qPCR on trauma and septic shock cohorts (Figures 2B, 5C). The designs targeting MLT1H or close neighborhood (PCR3, 4 and 5) presented the same profile, with a significant difference in septic shock and trauma cohorts compared to HV (PCR4). The design targeting the gene showed also up-modulation of CD55 and a very high absolute normalized expression in patients compared to HV (Figure 5C). [Of note 1555950_a_at probeset, targeting most of CD55 transcripts, was also up-modulated in patients, with a high expression level (data not shown)]. We also confirmed by flow cytometry on monocytes and neutrophils that CD55 expression was higher in patients than in HV, confirming an up-modulation at the protein level in patients (Figure 5D).

The MLT1D HERV element, targeted by the 1553043_a_at probe, is located in 3′ UTR of CD300LF-201, 202, 203, 204, and 207 protein-coding transcripts (Figure 6A). We made several RT-qPCR designs, targeting either the HERV locus only (PCR1)
or both HERV and 3′UTR of **CD300LF** (PCR2, Figure 6B). We confirmed the expression of the HERV locus, but the up-modulation seen in burn and septic shock 2 cohorts compared to HV on microarray was not confirmed by RT-qPCR, neither for gene nor for HERV designs (Figures 2C, 6C). PCR1 showed no signal at all. PCR2 design showed a slight higher expression level in burn and septic shock cohorts compared to HV. We also confirmed an higher expression at the protein level by flow cytometry on neutrophils in burn and septic shock patients, compared to HV (Figure 6D). In monocytes, protein level in burn patients at D1 seemed slightly higher than HV.

### DISCUSSION

We took advantage of previous microarray analyses on four cohorts of severely injured patients to assess the modulation of HERV transcriptome in acute inflammation. We showed that several loci were expressed and modulated after acute injury. Surprisingly, a large majority among the modulated HERVs were down-modulated in patients compared to HV, whereas a global and massive gene up-modulation has been observed after severe injuries (17).

Five HERVs were modulated in patients compared to HV in all four datasets and 16 HERVs in at least 3 datasets, suggesting a similar inflammatory triggered modulation in all models. We validated expression profiles by RT-qPCR on 6 regions, allowing us to explore more precisely the modulation pattern of the HERVs and the neighbor genes. Interestingly, all these 6 HERVs have detected signals in RNAseq experiments from lymphoid cells and whole blood datasets [Ensembl Rnaseq tracks (33)]. Some authors already focused on HERV detection in blood of septic patients (34, 35).

Moreover, very few data are available in human diseases for specific loci. No study had yet evaluated the expression of HERVs in acute inflammatory contexts by using multiple cohorts with different types of inflammatory injuries.

Several groups showed that huge epigenetic modifications occur after acute inflammation, regulating transcriptional profiles in the immune system, especially in sepsis (34, 35). These epigenetic modifications may explain the polarization profiles such as tolerance or trained immunity, observed after various stimulations of innate cells (35). We hypothesized and confirmed in vivo that other elements than genes, especially HERVs which are known to be tightly controlled by epigenetic modifications (36), might be modulated in acute inflammatory situations. This has also been demonstrated in other pathophysiological contexts such as cancer (7, 8, 37, 38), where global epigenetic modifications are also observed (39, 40).

Interestingly in cancer, epigenetic modifications that gave access to HERV cis sequences through open chromatin, have also revealed a very role in pathophysiology (2, 3, 37). Indeed, by providing alternative promoter sequences to classical protein coding genes, these epigenetic modifications explain part of the ectopic expression of myeloid-growth factor receptors in lymphoid cells (37). Recently, it has been suggested that HERV could provide transcription factor binding sites, modulating immune-related gene expression, as illustrated by MER41 elements which bring STAT1 motifs to IFNγ inducible genes (41). An exhaustive study on HERV expression with a different tool, like CHIPseq technology, would bring valuable data to find potential TFBS on HERV sequences. This underlines how HERV elements, in particular their LTRs, could modulate gene expression and the host immune response to injury. In our study, the four commonly modulated HERVs were LTRs located nearby genes related to the immune response. In several cases (**NFE4**, **CD55**, **CD300LF**, **SLC8A1**).
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CD300LF), we found a polyadenylation signal (AAUAAA) provided by the HERV LTR in 3' of some of the alternative transcripts of the genes. The case of CD300LF is interesting as this protein acts as an inhibitory receptor for myeloid cells (42). The LTR might stabilize specific transcripts and enhance expression of CD300LF protein, which we confirmed by flow cytometry in severe burn patients early after admission. This up regulation might participate in the compensatory anti-inflammatory response. The precise understanding of the mechanisms through which specific HERV LTRs might impact immune gene expression is not possible in such translational research setting with patient samples. This will require in the future in vitro experimental models to validate and understand our observations.

Our RT-qPCR validation assays also showed inter-individual variability and underlined that exploring such repertoire of our genome, repetitive sequences, may face specificity issues, and will require specific tools. Indeed, as a first attempt, we used commercial microarray where probesets were not initially designed to target HERV elements. Moreover, as the probesets targeting HERVs were initially supposed to target conventional genes, the majority of explored HERVs are close to or within a gene. To better understand HERV expression in these settings, targeting HERVs localized far from genes seems important. Until now, the lack of tool made difficult the exploration of HERV expression. It would be interesting to reproduce these analyses, with a more exhaustive technology designed to specifically target HERVs, like the HERV-V3 Affymetrix microarray we recently published (43), or even RNAseq. It will allow us to better describe the whole HERV transcriptome modulation and understand the putative global role of HERV in the host response.

Finally, it would be of importance to consider HERV expression in further blood transcriptome analyses, especially in such acute inflammatory contexts, to better understand HERV expression during host response. Such studies, based on well-defined cohorts including criteria for patient stratification and taking into account drug treatment, should allow to estimate whether HERVs could be good markers of the different immune phases that follow acute injury. More, whether HERV could become potential therapeutic targets would basically require to decipher which circulating cell type produces which HERV. It will thus be appropriate to develop dedicated cellular models to, in one hand better understand the contribution
of each blood-cell type to HERV expression and in another hand how HERV expression may contribute to the cell response to stimuli. To conclude, we showed for the first time that specific HERV loci are transcribed in whole blood of ICU patients. Our design allowed us to identify specific transcriptional signatures of HERV's elements, in vivo, linked
to the acute inflammatory response. Moreover, the similarities observed in three models of acute injuries suggest common regulatory mechanisms and a specificity of the observed modulation. We also unravel the potential regulatory role of these elements within the host immune response. Further studies are needed to better understand such mechanisms and how HERVs may contribute to the pathophysiology of the host immune response, a key part of the pathophysiology of sepsis.

**DATA AVAILABILITY STATEMENT**

Microarray expression data has been deposited on NCBI Gene Expression Omnibus and are accessible through GEO accession numbers GEO:GSE77791, GEO:GSE57065 and GEO:GSE95233. Data from microarray experiment for trauma cohort are available at Hospices Civils de Lyon—bioMérieux—UCBL1 "Pathophysiology of Injury Induced Immunosuppression," Groupement Hospitalier Edouard Herriot, France.
AUTHOR CONTRIBUTIONS

OT, FM and JT designed the project, performed the analyses and wrote the paper. CJ and FV performed cytometry experiments. EC performed RT-qPCR validations. ML, AL, BA, and TR recruited patients in the various cohorts. OT, MM, FV, FM, and JT read and discussed the manuscript. All authors drafted or revised critically the manuscript for important intellectual contents. All authors read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fimmu.2018.03091/full#supplementary-material

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Table S1 | Clinical description of trauma cohort.
Table S2 | Designs of qPCR systems. (A) Home-made designs. (B) Commercial designs.
Table S3 | Differentially expressed HERVs. The 56 differentially expressed probesets targeting HERVs between patients at D1 and HV, in at least 1 dataset. Each tab correspond to a dataset. Each table shows the probeset names, the log2 fold changes, the Average Expression, the moderated t-statistic, the raw p-value and the adjusted p-value.

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Conflict of Interest Statement: OT, MM, CJ, EC, AP, FM, and JT are employees of an in-vitro diagnostic company. The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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