Extracellular vesicles from human liver stem cells restore argininosuccinate synthase deficiency

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Abstract

Background: Argininosuccinate synthase (ASS1) is a urea cycle enzyme that catalyzes the conversion of citrulline and aspartate to argininosuccinate. Mutations in the ASS1 gene cause citrullinemia type I, a rare autosomal recessive disorder characterized by neonatal hyperammonemia, elevated citrulline levels, and early neonatal death. Treatment for this disease is currently restricted to liver transplantation; however, due to limited organ availability, substitute therapies are required. Recently, extracellular vesicles (EVs) have been reported to act as intercellular transporters carrying genetic information responsible for cell reprogramming. In previous studies, we isolated a population of stem cell-like cells known as human liver stem cells (HLSCs) from healthy liver tissue. Moreover, EVs derived from HLSCs were reported to exhibit regenerative effects on the liver parenchyma in models of acute liver injury. The aim of this study was to evaluate whether EVs derived from normal HLSCs restored ASS1 enzymatic activity and urea production in hepatocytes differentiated from HLSCs derived from a patient with type I citrullinemia.

Methods: HLSCs were isolated from the liver of a patient with type I citrullinemia (ASS1-HLSCs) and characterized by fluorescence-activated cell sorting (FACS), immunofluorescence, and DNA sequencing analysis. Furthermore, their differentiation capabilities in vitro were also assessed. Hepatocytes differentiated from ASS1-HLSCs were evaluated by the production of urea and ASS enzymatic activity. EVs derived from normal HLSCs were purified by differential ultracentrifugation followed by floating density gradient. The EV content was analyzed to identify the presence of ASS1 protein, mRNA, and ASS1 gene. In order to obtain ASS1-depleted EVs, a knockdown of the ASS1 gene in HLSCs was performed followed by EV isolation from these cells.

Results: Treating ASS1-HLSCs with EVs from HLSCs restored both ASS1 activity and urea production mainly through the transfer of ASS1 enzyme and mRNA. In fact, EVs from ASS1-knockdown HLSCs contained low amounts of ASS1 mRNA and protein, and were unable to restore urea production in hepatocytes differentiated from ASS1-HLSCs.

Conclusions: Collectively, these results suggest that EVs derived from normal HLSCs may compensate the loss of ASS1 enzyme activity in hepatocytes differentiated from ASS1-HLSCs.

Keywords: Liver transplantation, Citrullinemia type I, Stem cells, Extracellular vesicles

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Background

Citrullinemia type I is an autosomal recessive disorder caused by a deficiency in argininosuccinate synthase (ASS; common gene symbol alias: ASS1), an enzyme that catalyzes the third reaction of the urea cycle [1]. Typical symptoms that characterize the disorder include hyperammonemia, low plasma arginine levels, and elevated levels of citrulline in the plasma and urine [2, 3]. Furthermore, accumulation of ammonia during the first few days of life leads to poor feeding, vomiting, neuro-psychiatric symptoms, and coma [2]. Current treatments involve pharmacologic nitrogen scavenger therapy, lifelong protein restriction, and L-arginine supplementation which are aimed towards symptomatic control. For patients that do not respond satisfactorily to these treatments the only alternative is liver transplantation. However, due to the limited availability of donor livers, alternative treatments such as gene therapy are currently under investigation [2].

Previously, we have isolated a population of stem cell-like cells from the human liver known as human liver stem cells (HLSCs) which not only express liver tissue-specific markers, but also mesenchymal and embryonic stem cell markers [4, 5]. Furthermore, under appropriate cell culture conditions, HLSCs demonstrate in vitro differentiation abilities as well as in vivo regenerative properties [4, 5]. For instance, under low gravity culture conditions in the presence of hepatocyte growth factor and fibroblast growth factor 4, HLSCs differentiate into mature hepatocytes [4].

Over the past decade, stem cells have been studied extensively as a form of therapy due to their healing ability. Several preclinical models have been developed out of which a paracrine mechanism of action has been suggested [6, 7]. In the midst of various paracrine mediators identified, extracellular vesicles (EVs) have also been implicated to play a role in the exchange of information [8]. EVs are membrane bound particles secreted by cells in the extracellular milieu. They are classified into two main categories according to their size and subcellular origin. Exosomes are smaller in size (30–100 nm) and are released by exocytosis through the fusion of intracellular multivesicular bodies with the plasma membrane. Microvesicles, on the other hand, are larger (up to 1000 nm) and are shed by direct budding from the cell plasma membrane [8]. Once secreted, these vesicles either remain in the extracellular matrix close to the cell of origin or disperse to other parts of the body through the blood or lymphatic system allowing intercellular exchange of proteins, bioactive lipids, and nucleic acids.

As intercellular vehicles, EVs carry a variety of cargo. For instance, Ratajczak et al. demonstrated that EVs derived from embryonic stem cells may contribute towards the reprogramming of hematopoietic progenitor cells through the delivery of embryonic stem cell-derived proteins and mRNA [9]. Subsequently, other studies followed this up that showed that EVs derived from adult cells also contained and transferred miRNAs to target cells [10, 11].

The ability of EVs to reprogram as a consequence of transferring RNAs has been demonstrated in different tissues and pathologies [12]. For instance, we observed that endothelial progenitor cell-derived EVs activated angiogenic pathways in recipient endothelial cells through horizontal transfer of mRNA [11]. Moreover, in different models of tissue injury, treatments with EVs derived from mesenchymal stem cells have been reported to promote tissue regeneration [13–16]. We also observed that the administration of EVs derived from HLSCs improved the regeneration of liver in rats with a 70% hepatectomy [17] and assisted in the overall recovery of mice in a murine model of acute kidney injury [18].

In this study, we investigated whether EVs derived from normal HLSCs had the ability to carry and transfer wild-type ASS1 to mature hepatocytes differentiated from ASS1-mutated HLSCs, thereby restoring enzyme activity and urea production.

Methods

Isolation and culture of normal and mutated human liver stem cells

A liver specimen from a 5-year-old citrullinemia type I patient undergoing liver transplant was obtained under written informed consent from the parents. The tissue was provided anonymously according to the local tissue banking protocol of discarded tissues (# 80911) approved by the University of Torino Ethics Board (Comitato Bioetica di Ateneo).

Hepatocytes were isolated from the liver specimen (5–20 g) by collagenase digestion following the same protocol as for the isolation of normal HLSCs as described previously [5]. Briefly, isolated liver tissues were digested in liver digest medium (Invitrogen, Carlsbad, CA, USA) at 37 °C. The cell suspension obtained was filtered through a sterile 100-μm nylon mesh, centrifuged at 3000 g for 5 min and subsequently washed in cold wash medium (Invitrogen). The hepatocytes obtained were cultured in a medium composed of three parts α-minimum essential medium and one part endothelial cell basal medium-1 (3:1; Lonza, Basel, Switzerland) supplemented with l-glutamine (5 mM), penicillin (50 IU/ml), streptomycin (50 g/ml) (all from Sigma-Aldrich, St. Louis, MO, USA), and 10% fetal calf serum (FCS; Invitrogen). After a few days of culture ASS1-HLSC clones were expanded and subsequently frozen. Wild-type HLSCs were isolated from cryopreserved normal human hepatocytes (hH) obtained from Cambrex Bio Science, Verviers S.p.r.l. (Verviers, Belgium). hH were also used as normal control of mature hepatocytes.

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Isolation of HLSC-derived EVs (HLSC-EVs)
Briefly, 2 × 10^6 HLSCs and shRNA ASS1 knockdown HLSCs (HLSC shRNA-ASS1) were cultured in serum-free Roswell Park Memorial Institute (RPMI) medium for 18 h at 37 °C. The supernatant was centrifuged for 20 min at 3000 g to remove cell debris and apoptotic bodies. This was followed by a two-step ultracentrifugation protocol whereby the supernatant was first centrifuged at 10,000 g (10 K) for 1 h at 4 °C to pellet the 10 K EV fraction and then a second ultracentrifugation at 100,000 g (100 K) for 1 h at 4 °C to isolate the 100 K EV fraction (Beckman Coulter Optima L-90 K, Fullerton, CA, USA) [19]. Both the 10 K and 100 K fractions were resuspended in RPMI supplemented with 1% dimethyl sulfoxide (DMSO) and frozen at −80 °C for later use. The viability of HLSCs at the time of EV collection was 97–99% as determined by trypan blue exclusion.

As suggested by Kowal et al. [20], an iodixanol (Optiprep from Sigma-Aldrich) floating separation protocol was used to further purify the EV fractions from free floating contaminating proteins and RNAs [21]. Briefly, the pellets of EVs obtained after differential ultracentrifugation were directly resuspended in 500 μl 60% iodixanol (Optiprep from Sigma-Aldrich) mixed with 0.25 M sucrose and transferred to ultracentrifuge tubes (10.4-ml polycarbonate centrifuge bottles; Beckman Instruments). Subsequent gradients of iodixanol at 30%, 15%, and 5% were then layered on top of the initial 60% EV-iodixanol/sucrose preparation and the final volume adjusted to 10 ml by topping up with saline solution. The tubes were then ultracentrifuged at 350,000 g for 1 h, 4 °C, without braking in an Optima L-100 K ultracentrifuge (Beckman Coulter) equipped with type 90Ti rotor. Following ultracentrifugation, the 15%, 30%, and 60% fractions were recovered, diluted with phosphate-buffered saline (PBS), and ultracentrifuged again at 100,000 g for 1 h. The pellet obtained was resuspended in RPMI supplemented with 1% DMSO and either used fresh or stored for subsequent studies under −80 °C conditions. No difference in biological activity was observed between fresh and stored EVs.

In order to study the internalization of EVs by fluorescent microscopy, EVs were labeled with 1 μM Dil dye (Thermo Fisher Scientific, Waltham, MA, USA) as described previously [18]. Briefly, purified EVs were resuspended in PBS supplemented with 1 μM Dil dye and ultracentrifuged at 100,000 g for 1 h at 4 °C. Following labeling, the EVs were washed once more with PBS by ultracentrifugation as mentioned above. The pellet obtained was then resuspended in RPMI with 1% DMSO and frozen for subsequent studies.

Concentration and size distribution of EVs were determined by the Nanosight LM10 system (NanoSight, Wiltshire, UK). Briefly, EV preparations were diluted (1:200) in sterile saline solution and analyzed by the Nanoparticle Analysis System using the NTA 1.4 Analytical Software [21].

Flow cytometric analysis
Cytofluorometric analysis was performed as described previously [4]. The antibodies used (conjugated with phycoerythrin (PE) or fluorescein isothiocyanate (FITC)) are as follows: anti-CD45, anti-Albumin, anti-C90, anti-CD73, anti-CD105, anti-CD29, anti-CD31, anti-KDR, and anti-VE-Cadherin (Dako Denmark A/S, Copenhagen, Denmark). All incubations with antibodies were performed in 100 μl PBS containing 0.1% bovine serum albumin (BSA) at 4 °C. The cells were washed twice between incubations and analyzed (10,000 cells/sample) on a BD FACSCalibur cytometer (BD Biosciences Pharmingen, San Jose, CA, USA). For detection of albumin, cells were permeabilized and labeled with anti-albumin monoclonal antibody (R&D Systems, Abingdon, UK) followed by PE-conjugated anti-mouse IgG secondary antibody. All samples were gated on the basis of negative controls, and compensated appropriately prior to analyses. Population percentages and numbers were generated for gated populations from each experiment using Cell Quest software (BD Biosciences Pharmingen).

In vitro osteogenic, adipogenic, and endothelial differentiation
Osteogenic, adipogenic, and endothelial differentiation of ASS1-HLSCs was performed as described previously [4]. Briefly, ASS1-HLSCs were cultured in osteogenesis induction medium (Lonza) for 3 weeks (replenished twice per week for 3 weeks). Differentiation was evaluated by fixing cells with 4% paraformaldehyde for 20 min and then staining with alizarin red, pH 4.1 (Sigma-Aldrich), at room temperature. For adipogenic differentiation, cells were cultured in adipogenesis induction medium (Lonza), followed by adipogenic maintenance medium (Lonza). To evaluate the differentiation, cells were fixed with 4% paraformaldehyde for 20 min at room temperature then stained with 0.5% oil red O (Sigma-Aldrich) in methanol (Sigma-Aldrich). Endothelial cell differentiation was obtained by culturing ASS1-HLSCs in EBM (Lonza) with vascular endothelial growth factor (10 ng/ml; Sigma-Aldrich), at room temperature. For adipogenic differentiation, cells were cultured in adipogenesis induction medium (Lonza), followed by adipogenic maintenance medium (Lonza). To evaluate the differentiation, cells were fixed with 4% paraformaldehyde for 20 min at room temperature then stained with 0.5% oil red O (Sigma-Aldrich) in methanol (Sigma-Aldrich). Endothelial cell differentiation was obtained by culturing ASS1-HLSCs in EBM (Lonza) with vascular endothelial growth factor (10 ng/ml; Sigma-Aldrich) for 10 days. Following culture, endothelial differentiation was evaluated by flow cytometric analyses of endothelial markers as described previously [4].

Immunofluorescence analysis
Indirect immunofluorescence of cells was performed as described previously [4]. Briefly, ASS1-HLSCs cultured on chamber slides (Nalge Nunc International, Rochester, NY, USA) were fixed with 4% paraformaldehyde and/or permeabilized (only for intracellular markers) with 0.1%
Triton X-100 buffer. Cells were then labeled with the following monoclonal antibodies: anti-albumin (R&D Systems), anti-nestin (BD Biosciences Pharmingen), anti-α-fetoprotein (aFP; R&D Systems), anti-nanog (Abcam, Cambridge, MA), anti-Sox2 (Abcam), anti-vimentin (Sigma-Aldrich), anti-cytokeratin 8 (CK8), anti-CK19, anti-Von Willebrand factor (all from Sigma-Aldrich), and anti-ASS1 (Thermo Fisher Scientific). Alexa Fluor 488 anti-rabbit IgG and Texas Red anti-mouse IgG (Thermo Fisher Scientific) were used as secondary antibodies and Hoechst 33258 dye (Sigma-Aldrich) was applied for nuclear staining. Labeling of cells with only secondary antibodies or substitution with nonimmune rabbit, rat, or mouse IgGs served as controls. Slides were analyzed by confocal microscopy using a Zeiss LSM 5 Pascal Model Confocal Microscope (Carl Zeiss International, Jena, Germany).

**In vitro culture in RCCS and urea production**

In order to differentiate HLSCs and ASS1-HLSCs into functional hepatocytes, cells were cultured in microgravity conditions using the Rotary Cell Culture System (RCCS; Synthecon Incorporated, Houston, TX, USA) as described previously [4]. Briefly, cells (250,000/ml) were resuspended in medium consisting of 75% Dulbecco's modified Eagle's medium (DMEM; Mediatech, Herndon, VA, USA), and 25% MCDB 201 (Sigma-Aldrich) supplemented with 2% fetal bovine serum (FBS), linoleic acid (Sigma-Aldrich), L-ascorbic acid (Sigma-Aldrich), insulin-transferrin-selenium (ITS; Sigma-Aldrich), penicillin (100 μg/ml), streptomycin (100 μg/ml), and 10 ng/ml of hepatocyte growth factor and fibroblast growth factor 4; 10 ml of the cell suspension was then loaded into circular RCCS vessels and rotated about their own axis at 8–10 rotations per minute under 37 °C conditions. After 4 days of culture, the supernatant was collected, centrifuged at 300 g for 5 min and stored at –20 °C. Levels of urea in the supernatants were then evaluated using the blood urea nitrogen (BUN) colorimetric detection kit as per the manufacturer's protocol (Arbor assays, MI, USA). For selective RCCS experiments, cells were cultured in the presence of HLSC-EVs with or without α-methyl-DL-aspartic acid (MDLA), a specific inhibitor of ASS1 enzyme. Treatment of ASS1-HLSCs with ASS1 shRNA EVs or fibroblast EVs (hFib EV) served as controls.

**Western blot analysis**

The level of ASS1 enzyme in cells was determined by Western blot analyses. Briefly, cells were lysed on ice for 30 min in radioimmunoprecipitation assay (RIPA) buffer (50 mM Tris-HCl PH 7.4, 150 mM NaCl, 1% NP-40, 0.1% SDS; Sigma-Aldrich) supplemented with the protease inhibitors phenylmethylsulfonyl fluoride (PMSF) (1 mM), leupeptin (10 μg/ml), and aprotinin (100 units/ml). Proteins from EVs were isolated using the all-in-one purification kit (Norgen Biotek Corp, TO, Canada) according to the manufacturer’s protocol, and the concentrations determined by the Bradford method.

Protein samples at a concentration of 10–30 μg were separated in 8% or 4–15% gradient SDS-PAGE gels under reducing conditions and electroblotted onto 0.2-mm nitrocellulose membranes (GE Healthcare Life Sciences, MA, USA). The membranes were blocked in Tris-buffered saline-Tween (TBS-T; 25 mM Tris, pH 8.0, 150 mM NaCl, and 0.05% Tween-20) containing 5% (w/v) nonfat dried milk for 1 h. After blocking, membranes were probed overnight with mouse anti-ASS1 (BD Transduction Laboratories), mouse anti-CD63, rabbit anti-tubulin, and goat anti-actin (Santa Cruz Biotechnology, CA, USA). After extensive washings with TBS-T, the blots were incubated with appropriate peroxidase conjugated secondary antibodies (goat anti-mouse, goat anti-rabbit, and mouse anti-goat IgG; Pierce, Thermo Fisher Scientific) for 1 h at room temperature. Following incubation, the membranes were washed extensively with TBS-T, probed with enhanced Super Signal West Femto Maximum Sensitivity Substrate (Thermo Fisher Scientific), and detected by the Chemidoc system (Bio-rad, CA, USA).

**Immunoprecipitation and enzymatic activity of ASS1**

Cells were lysed in non-denaturing lysis buffer (137 mM NaCl, 2 mM EDTA, 20 mM Tris–HCl, pH 7.5, 10% glycerol, 1% Nonidet P-40, and protease inhibitor cocktail) and centrifuged at 300 g for 15 min at 4 °C. The supernatants were collected and quantified for total protein concentration using the Bio-Rad protein assay method.

Immunoprecipitation of ASS1 was performed using Dynabeads Protein G (Thermo Fisher Scientific). Briefly, the beads were incubated with 10 μg ASS1-antibody (Thermo Fisher Scientific) per mg of total protein for 1.5 h at 4 °C to form complexes. The bead/ASS1-antibody complexes were washed with PBS-0.02% tween and incubated with cell lysates (1 mg total protein) overnight at 4 °C under rotation. After three washes with PBS-0.02% tween, ASS1 proteins were eluted by adding 20 μl 50 mM glycine at pH 2.8. In order to prevent the loss of the tridimensional structure of ASS1, 200 μl TRIS-HCL 20 mM pH 7.8 was added.

Eluted proteins were then evaluated for ASS1 activity using a modified version of the assay described by Lakhal-Naouar et al. [22]. Briefly, eluted protein samples (12.5 μl per reaction) were resuspended in reaction buffer (20 mM Tris-HCl, pH 7.8, 4 mM ATP, 4 mM citruline, 4 mM aspartate, 6 mM MgCl2, 20 mM KCl, and 0.2 units of pyrophosphatase) to a final volume of 20 μl. The reaction samples were then transferred to 96-well microtiter plates in duplicates and incubated for 30 min at 37 °C. Following incubation the reaction was stopped by adding 20 μl malachite green reagent (Malachite
Green Phosphate Assay Kit, Bioassay System, CA, USA). Reactions without citrulline and aspartate substrates were prepared to serve as controls. Phosphate accumulation was determined at 655 nm by spectrophotometry, and the concentration interpolated from a standard curve of inorganic phosphate (Pi). Due to the spontaneous release of Pi in the absence of substrates, the final concentration of Pi from affinity-purified proteins and/or rat liver extract (RLE; served as positive control for the reaction) were determined by subtracting the mean concentration of Pi obtained in the absence of substrates from the mean concentration of Pi obtained in the presence of substrates. The specific ASS1 activity was determined using the following formula: nanomoles of phosphates released/mg protein/30 min [22]. In selected experiments, ASS1-siRNA transfected ASS1-HLSC were used.

RNA extraction and qRT-PCR
Total RNA from cells was extracted using TRIzol reagent (Thermo Fisher Scientific) and total RNA from HLSC-EVs was isolated using the all-in-one purification kit (Norgen Biotek Corp.) according to the manufacturer’s protocol. The quantity and quality of isolated RNAs were determined using the Nanodrop ND-1000 spectrophotometer (Thermo Fisher Scientific). cDNA was then synthesized using the High Capacity cDNA Reverse Transcription Kit (Thermo Fisher Scientific) as per the manufacturer’s protocol. cDNA at a concentration of 2 ng was used per sample per quantitative real-time polymerase chain reaction (qRT-PCR) reaction in triplicate using a 96-well StepOne Real-Time System (Thermo Fisher Scientific). RNAs were determined using the Nanodrop ND-1000 spectrophotometer (Thermo Fisher Scientific). cDNA was then synthesized using the High Capacity cDNA Reverse Transcription Kit (Thermo Fisher Scientific) as per the manufacturer’s protocol. cDNA at a concentration of 2 ng was used per sample per quantitative real-time polymerase chain reaction (qRT-PCR) reaction in triplicate using a 96-well StepOne Real-Time System (Thermo Fisher Scientific). Wells containing only primers were cycled in parallel with each run and served as negative cDNA controls. For mRNA expression in cells, results were analyzed using the RQ method with GUSB gene as an internal control and for HLSC-EVs, the 2−ΔΔCt method was adopted using S18 gene as internal control. Primers and probes were designed using the Universal ProbeLibrary Assay Design Center software (www.lifescience.roche.com). Primers used to amplify ASS1 mRNAs are summarized in Table 1.

ASS1 silencing
A set of five pLKO.1 HIV-based lentiviral vectors targeting the human ASS1 gene were purchased (TRCN0000045553, TRCN0000045554, TRCN0000045555, TRCN0000045556, TRCN0000045557; Dharmacon RNAi Consortium (TRC) Lentiviral shRNA). shRNAs were tested on HLSCs to evaluate the silencing efficiency and the most efficient and stable shRNA was used to specifically downregulate ASS1 expression in HLSCs. The lentiviruses pLKO.1-scr (expressing a “scramble” shRNA as control) and pLKO.1-ASS1 (expressing the shRNA specific for ASS1) were produced in HEK293FT cells using a lentiviral packaging system after which HLSCs were infected with the lentiviruses in the presence of Sequa-brene (Sigma-Aldrich). Following infection, cells were selected with 5 μg/ml puromycin. The efficacy and stability of silencing in selected cells was evaluated by qRT-PCR.

siRNA transfection
ASS-siRNA and nonsilencing siRNA were purchased from Thermo Fisher Scientific. Transient transfection of ASS-siRNA was performed using Lipofectamine RNAiMAX transfection reagent (Invitrogen) as per the manufacturer’s protocol. ASS1-HLSCs (500,000 cells per well) were seeded in a petri dish (10-cm diameter) and cultured for 24 h. The complete culture medium was replaced with antibiotic-free medium and transfected with 300 pM ASS-siRNA or nonsilencing RNA for 3 days in the presence of Lipofectamine RNAiMAX. Before the addition in cell cultures, siRNAs were incubated with lipofectamine for 20 min at room temperature in Opti-Mem medium (Invitrogen). Preliminary experiments using FITC-siRNA (Qiagen, Valencia, CA, USA) indicated that the peak of expression was 3 days after transfection. After 3 days of transfection, ASS1-HLSCs were detached with trypsin and 3.0 × 106 cells were cultured in RCCSs in the presence of 1 × 1010 EV-HLSCs. The effective ASS-siRNA transfection was evaluated in ASS1-HLSCs by RT-PCR. ASS enzymatic activity was measured after 4 days of culture under RCCS culture conditions.

DNA mutation analysis
Genomic DNA was extracted using DNeasy Blood & Tissue Kit (Qiagen) according to the manufacturer’s instructions. The quality and quantity of extracted DNA was determined using the Nanodrop ND-1000 spectrophotometer (NanoDrop Technology).
Two regions of the human ASS1 gene located in exons 14 and 15 were amplified by PCR. These regions comprise of the following codon mutations: R363W (exon 14) and G390R (exon 15). The primers were developed (Table 2) based on the GenBank reference sequence (accession no. NC_000009.11), using Primer Express 3.0 software (Thermo Fisher Scientific). PCR was then performed in a final volume of 50 μl, containing 10× PCR buffer, 0.05 units/μl JumpStart Taq polymerase, 10 mM deoxynucleotide triphosphates mix, 150 ng genomic DNA, and 500 nM primers. Cycling conditions were as follows: 1 min at 94 °C, 35 cycles at: 94 °C for 30 s, 62 °C

| SNaPshot primers | SNaPshot primer sequences |
|------------------|---------------------------|
| R363W            | T26GGTGTAACCTCTCGGC       |
| G390R            | T35GCCAACGTATGCGC         |

R363W was used to amplify the genomic region containing the codon mutation g.55277 C > T and primer G390R for the region containing the mutation g.59839 G > A.

Fig. 1 Characterization of ASS1-HLSCs. 

**A** Representative flow cytometric analysis of markers expressed by ASS1-HLSC (black histograms; dotted histograms represent the isotypic controls). Cells were labeled with FITC- or PE-conjugated antibody. One hundred percent of ASS1-HLSCs were positive for CD90, CD73, CD29, and CD105, and negative for CD45 as described previously for normal HLSCs [4]. ASS1-HLSCs were also positive for the liver marker albumin [4]. Data represent one of three experiments with similar results. 

**B** Representative immunofluorescence micrographs of ASS1-HLSCs stained with antibodies against albumin, nestin, α-fetoprotein (α-FP), nanog, sox2, vimentin, cytokeratin (CK)8 and CK19. Hoechst dye 33258 was applied for nuclei staining. Scale bars = 5 μm. Data represent one of three experiments with similar results.
for 30 s, and 72 °C for 60 s, followed by 1 min at 72 °C. At the end of the reaction, 10 μl PCR-amplified DNA samples were analyzed using 1% agarose gel.

To evaluate the presence of DNA in EVs derived from HLSCs, DNA was extracted and analyzed by PCR. Briefly, 14 different preparations of EVs obtained from HLSCs were pooled together by ultracentrifugation at 100,000 g for 2 h at 4 °C. The supernatant was discarded and lysis buffer was added directly to the EV pellet and processed as per the manufacturer’s protocol to isolate the DNA (DNeasy Blood & Tissue Kit, Qiagen). Exons 14 and 15 of the hASS1 gene were amplified using PCR as described above and amplicons were analyzed using 1% agarose gel.

PCR products from ASS1-HLSCs and ASS1-HLSCs treated with HLSC-EVs were analyzed for mutations (c.1087C > T and c.168G > A) using the ABI PRISM SNaPshot Multiplex Kit (Thermo Fisher Scientific), according to the protocol supplied by the manufacturer (normal HLSCs served as control). The SNaPshot method is based on the dideoxy single-base extension of unlabeled oligonucleotide primers. For each of the two mutations analyzed above, a primer annealing adjacent to the potentially mutant nucleotide was developed (Table 2). Briefly, the SNaPshot reaction was performed in a volume of 10 μl, containing 3 μl PCR product, 5 μl ready reaction mix, 1x sequencing buffer, and SNaPshot primers concentrated at 0.02 pmol/μl. At the end of the reaction, the labeled products were first treated with shrimp alkaline phosphatase (SAP; USB Corporation) to remove excess dideoxynucleotide triphosphate and then separated on 36-cm-long capillaries by performing a 25-min run in an automatic sequencer (ABI PRISM 3500 Genetic Analyzer, Thermo Fisher Scientific). GeneScan Analysis Software version 3.7 (Thermo Fisher Scientific) was used for data analysis. For each sample, the intensity of the fluorescent peak was measured.

**Fig. 2** The expression of ASS1 gene and ASS1 mRNA in HLSCs and ASS1-HLSC. a, b Representative SNaPshot sequences of ASS1 gene in normal human liver stem cells (HLSCs) and HLSCs from the liver of a patient suffering from citrullinemia type 1 (ASS1-HLSCs). a The upper electropherogram shows a normal ASS1 gene profile from HLSCs; the lower electropherogram shows the mutated gene profile from ASS1-HLSCs (c.1087C > T). b The upper electropherogram shows a normal ASS1 gene profile; the lower electropherogram shows the mutated ASS1-HLSC gene profile (c.168G > A). Data represent one of three experiments with similar results. c qRT-PCR analysis showing the expression of ASS1 isoforms 1 (Iso1) and 2 (Iso2) mRNA in human hepatocytes (hH), HLSCs, and ASS1-HLSCs. hH were used as positive control. d qRT-PCR analysis of ASS1 Iso1 mRNA in HLSCs transfected with ASS1 shRNA. Data represent the mean of two independent experiments performed in triplicate. *p < 0.05, HLSC shRNA-ASS1 vs HLSC Ctrl-SCR. A adenine, C cytosine, F.U fluorescence unit, G guanine, SCR scramble, T thymine
**Statistical analysis**
Results are expressed as mean ± standard deviation (SD). Statistical analysis was performed using the Student’s t test and analysis of variance (ANOVA) with Newman-Keuls test or ANOVA with Dunnet’s multicomparison tests where appropriate. A p value <0.05 was considered significant.

**Results**

**Characterization and in vitro differentiation of citrullinemia type I-derived ASS1-HLSCs**

ASS1-HLSCs derived from the liver of a patient with citrullinemia type I were characterized to confirm their expression of stem cell markers similar to HLSCs. Flow cytometric analysis revealed that ASS1-HLSCs, like their normal counterparts, were positive for several mesenchymal stem cell markers such as CD90, CD73, CD29, and CD105 (Fig. 1a). Furthermore, they were positive for the liver tissue-specific marker albumin but negative for the hematopoietic marker CD45 (Fig. 1a). In addition, immunofluorescence microscopy confirmed ASS1-HLSCs to be positive for the hepatocyte precursor marker α-FP, as well as vimentin (100%), nestin (100%), and CK8 (40%) (Fig. 1b). In contrast, CK19, a marker for oval cells, was negative (Fig. 1b). To assess the in vitro differentiation capabilities of ASS1-HLSCs, they were induced to differentiate into endothelial, osteogenic, and adipogenic lineages. Similar to HLSCs, ASS1-HLSCs were able to undergo endothelial differentiation when cultured in endothelial induction medium (Additional file 1: Figure S1). After 3 weeks in culture, ASS1-HLSCs compared to undifferentiated cells expressed de novo endothelial markers such as KDR, CD31, VE-cadherin, and von Willebrand Factor (vWF). Furthermore, when cultured in osteogenic differentiation medium, ASS1-HLSCs transformed morphologically from spindle shaped cells to cuboidal cells and underwent mineralization as reflected by alizarin staining (Additional file 1: Figure S1). ASS1-HLSCs when maintained in adipogenic differentiation medium did not differentiate into adipocytes as was also observed with normal HLSCs (data not shown).

In order to identify and characterize the ASS1 mutations in ASS1-HLSCs, a SNaPshot sequencing analysis was implemented. Briefly, after DNA extraction, a first amplification step was performed using two different specific primers for the genomic regions of the ASS1 gene expressing two potential codon mutations (g.55277 C > T, g.59839 G > A). The PCR products were then analyzed for mutations using the ABI PRISM SNaPshot Multiplex Kit. As shown in Fig. 2a and b, the mutations in ASS1-HLSCs were confirmed by the presence of the mutated bases T and A which were absent in normal HLSCs. At variance to mature hH that expressed both isoform 1 and 2 of ASS1

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**Fig. 3** The expression of ASS1 protein in HLSCs and ASS1-HLSCs. **a** Western blot confirming the presence or absence of ASS1 protein isoform 1 (Iso1) and isoform 2 (Iso2) in 10 μg total protein extracted from human hepatocytes (hH), human liver stem cells (HLSCs), HLSCs from the liver of a patient suffering from citrullinemia type 1 (ASS1-HLSCs), and Rotary Cell Culture System (RCCS)-differentiated HLSCs and ASS1-HLSCs. Actin was run on a parallel gel as a representative of loading. **b** Quantification of the relative band intensity of ASS1 Iso1 across all samples in Fig. 3a normalized to actin. **c** Representative immunofluorescence micrographs of hH, HLSCs, ASS1-HLSCs, and RCCS-differentiated HLSCs and ASS1-HLSCs stained for ASS1 enzyme. The nuclei of the cells were stained with Hoechst dye 33258. Original magnification × 400, scale bar = 50 μm. Data represent one of two experiments with similar results. **d** Quantification of the relative fluorescence intensity (RFI) of all the cell culture conditions in Fig. 3c with hH serving as a positive control established as the 100%. Data represent the mean of three independent experiments with similar results. §p < 0.05, HLSCs/RCCS vs HLSC; ^p < 0.05, ASS1-HLSC/RCCS vs ASS1-HLSC; *p < 0.05, HSC/RCCS vs HLSC; #p < 0.05, ASS1-HLSC/RCCS vs ASS1-HLSC
mRNA, the undifferentiated HLSCs and ASS1-HLSCs expressed mainly isoform 1 (Fig. 2c). Figure 2d shows the reduction of ASS1 mRNA expression in HLSCs silenced for the ASS1 gene.

Expression of ASS1 protein
The expression of ASS1 protein in mature hH, HLSCs, and ASS1-HLSCs was evaluated by Western blot (Fig. 3a and b). hH expressed two isoforms of ASS1 protein, one
at 46 kDa (isoform 1 (Iso1)) and the other at 40 kDa (isoform 2 (Iso2)) (Fig. 3a). Consistent with the low expression of Iso2 mRNA in undifferentiated HLSCs and ASS1-HLSCs, Western blot analysis showed only Iso1. This suggests that under undifferentiating conditions the low amount of Iso2 mRNA was not translated into protein. However, differentiation into hepatocytes via RCCS induced the expression of the Iso2 for both HLSCs and ASS1-HLSCs (Fig. 3a). Moreover, after differentiation the expression of Iso1 was increased in both HLSCs and ASS1-HLSCs compared to the undifferentiated counterparts (Fig. 3a and b).

The expression of the ASS1 protein in hH, HLSCs, and ASS1-HLSCs was further confirmed by immunofluorescence staining (Fig. 3c). As seen in Fig. 3d, quantitative analysis of ASS1 protein expression shows a significant increase after RCCS-induced differentiation (Fig. 3c and d). These data therefore confirm that differentiation to hepatocytes of both the wild-type and mutated HLSCs significantly increases the expression of the ASS1 protein.

Characterization of HLSC-EVs

On analyzing the 10 K and 100 K fractions, ASS1 protein expression was observed only in the 100 K fraction concurrently with the expression of the exosomal marker CD63 (Fig. 4a). Furthermore, the same fraction was also positive for the mesenchymal markers typical of HLSCs as observed by fluorescence-activated cell sorting (FACS) analysis (Fig. 4b). No further analysis was performed on the 10 K fraction as it was negative for ASS1 and CD63 (Fig. 4a).

The 100 K fraction was also submitted to floating separation whereby 75% of the EVs were detected in the 15% iodixanol fraction with a mean size of 155 ± 3.6 nm as determined by NanoSight analysis (Fig. 4c and d). In addition, transmission electron microscopy revealed a homogenous layer of these vesicles with a morphology resembling exosomes with a smaller size than observed by NanoSight (Fig. 4c, insert). Furthermore, on analyzing the EVs obtained from the three iodixanol fractions by Western blot, only the 15% fraction was positive for the exosomal marker CD63 as well as the ASS1 protein (Fig. 4e).

PCR analysis also confirmed at a molecular level the presence of mRNA for ASS1 Iso1 but not for ASS1 Iso2 in 100 K HLSC-EVs (Fig. 4f) and in the 15% iodixanol EV fraction compared to the other fractions (Fig. 4g).

In order to evaluate whether restoration of ASS1 enzymatic activity was observed in ASS1-HLSCs treated with HLSC-EVs, we generated EVs depleted of the ASS1 enzyme by silencing the gene at a cellular level. As shown in Fig. 4h, EVs derived from HLSC shRNA-ASS1 were depleted of both Iso1 ASS1 mRNA and protein.

EVs derived from normal HLSCs increased urea production and ASS1 enzymatic activity in mutated ASS1-HLSCs

As shown in Fig. 5, HLSC-EVs were internalized in ASS1-HLSCs after 6 h of incubation at 37 °C. It has been described that, as a consequence of the genetic mutations, the ASS1 enzyme activity is decreased in

![Fig. 5](image-url)
HLSCs. HLSC-EV treatment significantly increased urea production and enzymatic activity in ASS1-HLSCs, ASS1-HLSCs were treated with HLSC-EVs (1 × 10¹⁰ particles) in RCCS for 4 days. At the end of the experiments, urea production and ASS enzymatic activity were measured (Fig. 6a and b). Mutated ASS1-HLSCs produced significantly less urea and enzymatic activity than normal ASS1-HLSCs. ASS1-HLSCs treated with EVs contained ASS1 protein and a low level of ASS1 mRNA (data not shown) were ineffective (Fig. 6a). Moreover, the reduction of ASS1 urea production by EVs derived from hFib also depended on an inhibition of cell differentiation in RCCS (data not shown).

**HLSC-EV treatment decreased DNA mutations in ASS1-HLSCs**

Since HLSC-EVs carry fragments of ASS1 DNA (Fig. 7), we evaluated whether they may affect the mutations when incorporated into ASS1-HLSCs. We performed SNaPshot sequencing to determine the contribution of HLSC-EVs to correction of both mutations present in ASS1-HLSCs. Figure 8 shows that the amplitude of picks of the mutated bases T and A belonging to one allele of the ASS1 region of ASS1-HLSCs was significantly reduced when cultures were cultured in adhesion culture for 6, 24, and 48 h with HLSC-EVs (1 × 10¹⁰ particles).

**Discussion**

In the present study we demonstrated that EVs derived from normal HLSCs increased the production of urea and ASS1 enzymatic activity in hepatocytes differentiated from HLSCs derived from a patient with Citrullinemia type I.

Citrullinemia type I is an inherited disorder of the urea cycle caused by a genetic mutation in the ASS1 gene. The absence of a functional form of ASS1 enzyme leads to a rapid accumulation of ammonia and other toxic substances in the blood leading to an early neonatal death. Liver transplantation remains the only curative treatment for this defect. However, due to limited availability of donor livers, substitute therapies are currently being researched [23].

Recently, Syres et al. demonstrated that the accumulation of cysteine in organs was decreased after the injection of mesenchymal stem cells in a cysteinotic knockout mice model (CTNA⁻/⁻) [24]. The cells were found in multiple organs resulting in a 90% decrease in tissue cysteine levels, leading to the recovery of kidney injury [24]. Nevertheless, the reduction in tissue cysteine levels could not be attributed to mesenchymal stem cell transdifferentiation or fusion with resident CTNA mutated cells. In further studies, to explain the mechanism...
of reduction of intralysosomal accumulation of cysteine, Iglesias et al. used an in vitro coculture model of CTNA–/– fibroblasts and human amniotic mesenchymal stem cells [25]. They demonstrated that the uptake of mesenchymal stem cell-derived EVs carrying wild-type cysteinosin protein and mRNA by mutant fibroblasts significantly reduced the lysosomal accumulation of cysteine in the cells. Katsuda et al. recently showed that EVs derived from adipose mesenchymal stem cells may transfer a functional enzyme to target cells [26]. In addition, several studies provided evidence of effective transfer of functional mRNA, miRNA, and DNA by EVs [27–29].

Collectively, these works suggest that EVs could interact with target cells and transfer functional protein and nucleic acids that may modify their phenotype. In agreement with this evidence, we found that HLSC-EVs contained wild-type ASS1 protein, mRNA, and DNA. Furthermore, on incubating these EVs with mutated ASS1-HLSCs the mutated enzymatic effect was corrected, restoring urea production and enzymatic activity. Undifferentiated HLSCs expressed Iso1 and, to a significantly less extent, Iso2 mRNA, and the amount of Iso2 protein was undetectable by Western blot analysis. Consistently, EVs secreted from undifferentiated HLSCs contained only mRNA and protein of Iso1. In addition, knocking down Iso1 by shRNA in HLSCs depleted the levels of ASS1 mRNA and protein in the EVs released subsequently by the cells. These EVs from ASS1 shRNA knockdown HLSCs were unable to restore urea production in hepatocytes differentiated from ASS1-HLSCs. This result suggests that the biological activity of EVs may depend on the transfer of both functional ASS1 protein and mRNA. It is unclear from this experiment to what extent protein/mRNA transfer contributed to the recovery of ASS1 enzymatic activity in ASS1-HLSCs. However, the result of the experiment using transfection of ASS1-HLSCs with ASS1-siRNA that shows inhibition of HLSC-EV-induced correction of enzymatic activity suggests a predominant role of mRNA transfer.

In addition, we also found that HLSC-EVs carried fragments of the nonmutated ASS1 gene and that the amplitude of the peak of the bases mutated was reduced in ASS1-HLSCs after treatment with vesicles. However, the transfer of DNA was not relevant for the correction of the enzyme activity in ASS1-HLSCs as EVs derived from HLSCs silenced with ASS1 shRNA (which only diminished ASS1 mRNA levels) had a significant reduction in their biological activity.

Further studies are required to evaluate whether the horizontal transfer of DNA could also correct the enzymatic defect in the mutated cells. Nevertheless, several studies have recently investigated the possibility of horizontal transfer of DNA by EVs. For instance, Lázaro-Ibáñez et al. [30] reported the presence of double-stranded genomic DNA fragments in exosomes derived from prostate cancer cells and suggested the transfer of oncogenes between cancer and noncancerous cells. Fischer et al. demonstrated the horizontal transfer of DNA genes by EVs in recipient cells [31]. These studies suggest that vesicles containing genomic DNA may be instrumental in the exchange of genetic information between eukaryotic cells and open the possibility to exploit this mechanism for correction of genetic disorders.

The mechanisms of exosomal biogenesis and sorting of proteins and nucleic acids are only partially known. However, recently the mechanisms controlling specific miRNA sorting in hepatocyte exosomes has been described by Santangelo et al. [32]. They identified an RNA binding protein SYNCRIp as a critical component of miRNA accumulation within exosomes, suggesting a
possible selective modulation of exosome cargo. EVs are enriched in ribonucleoproteins that may be involved in the storage, traffic, and protection of degradation of ribonucleic acid [33]. Moreover, the association of Alix, a component of endosomal sorting complex required for transport, with Ago2, which is involved in miRNA maturation, has been shown to contribute to miRNA accumulation into EVs [34].

Conclusions
EVs derived from stem cells may act as paracrine mediators by transferring proteins and nucleic acids inducing epigenetic changes in recipient cells. Herein, we found that EVs derived from normal HLSCs have the potential to correct ASS1 enzyme deficiency in hepatocytes differentiated from HLSCs derived from a patient with citrullinemia type I, suggesting a potential application of EVs derived from stem cells in some inherited diseases.

Additional file

Additional file 1: Figure S1. In vitro differentiation of ASS1-HLSCs. (A) Representative flow cytometric analysis of undifferentiated ASS1-HLSCs showing negative expression of KDR, CD31, and VE-Cadherin. Von Willebrand factor (vWF) expression and alizarin red staining for mineralization was also
negative in undifferentiated conditions as observed by microscopy. (B) Representative flow cytometric analysis of ASS1-HLSCs after endothelial differentiation showing the presence of KDR, CD31, and VE-Cadherin (black histograms; dotted histograms represent isotypic controls). Representative micrographs showing the expression of VWF in ASS1-HLSCs after endothelial differentiation, and osteogenic differentiation of ASS1-HLSCs showing positive staining for calcium deposits as indicated by alizarin 21 days after culturing in osteogenic differentiation medium. Scale bar = 50 μm. Data represent one of three experiments performed with similar results. (TIF 8299 kb)

Abbreviations
ASS: Argininosuccinate synthase; ASS1-HLSC: Human liver stem cell from the liver of a patient suffering from citrullinemia type I; Ck: Cytokeratin; DMSO: Dimethyl sulfoxide; EV: Extracellular vesicle; FACS: Fluorescence-activated cell sorting; FITC: Fluorescein isothiocyanate; Fix: human fibroblasts; hH: Human hepatocytes; HLSC: Human liver stem cell; Isc: Isosmotic; MDLA: α-Methyl-DL-aspartic acid; PBS: Phosphate-buffered saline; PE: Phycoerythrin; Pi: Inorganic phosphate; qRT-PCR: Quantitative real-time polymerase chain reaction; RCCS: Rotary Cell Culture System; RPMI: Roswell Park Memorial Institute; aFp: α-Fetoprotein

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Availability of data and materials
All data generated or analyzed during the current study are included in this article.

Authors’ contributions
MBHS and SP isolated ASS1-HLSCs, performed hepatocyte differentiation and ASS1 enzymatic activity assays, and contributed towards manuscript writing. SB and VF analyzed data and contributed towards manuscript writing. MCD performed Western blot analysis and iodixanol floating separation protocol. SK performed experiments, analyzed data, and contributed towards manuscript writing. SP performed ASS1 silencing experiments. ER contributed to the analysis and interpretation of data. RC performed DNA mutation analysis. RC, MSp, and MSa performed the metabolic studies, collected the liver specimen from the Liver Transplantation Centre, and contributed towards data assembly for the study. CT analyzed and interpreted data. GC contributed towards the conception, design, writing, and financial support. All authors read and approved the final manuscript.

Ethics approval and consent to participate
Human tissue was obtained from the native liver of a 5-year-old patient with neonatal-onset citrullinemia type I undergoing a liver transplant with the written informed consent of the parents. The liver specimen was acquired from the native liver removed for transplantation by the Liver Transplantation Centre according to a local tissue banking protocol (# 80911) approved by the University of Torino Ethics Board (Comitato Bioetica di Ateneo) and Centre according to a local tissue banking protocol (# 80911) approved by the University of Torino Ethics Board (Comitato Bioetica di Ateneo) and Centre according to a local tissue banking protocol (# 80911) approved by the University of Torino Ethics Board (Comitato Bioetica di Ateneo) and Centre according to a local tissue banking protocol (# 80911) approved by the University of Torino Ethics Board (Comitato Bioetica di Ateneo). Consent for publication
Not applicable.

Competing interests
MBHS, MCD, and GC are named as inventors in related patent applications. The remaining authors declare that they have no competing interests.

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