Effect of nutritional supplement on bone marrow-derived mesenchymal stem cells from aplastic anaemia

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Abstract

Aplastic anaemia (AA) is characterised by pancytopenia resulting from a marked reduction in haemopoietic stem cells (HSC). The regulation of haemopoiesis depends on the interaction between HSC and various cells of the bone marrow (BM) microenvironment, including BM-derived mesenchymal stromal cells (BMSC). The purpose of this study was to analyse the biological effect of nutritional supplement (NS), a dietary supplement consisting of thirty-six compounds: amino acids, nucleotides, vitamins and micronutrients on the BMSC of AA rats. The AA rat model was established by irradiating X-ray (2.5 Gy) and intraperitoneal injections of cyclophosphamide (35 mg/kg; Sigma) and chloramphenicol (35 mg/kg; Sigma). Then AA rats were fed with NS in a dose-dependent manner (2266, 95, 1511 mg/kg d) by intragastric administration. The effect of NS on the BMSC of AA rats was analysed. As compared with AA rats, NS treatment significantly improved these peripheral blood parameters and stimulated the proliferation of total femoral nucleated cells. NS treatment affected osteogenic differentiation of BMSC and enhanced bone mineral density. Co-incubation of HSC with mesenchymal stromal cells and serum from AA rats subjected to high-dose NS markedly improved the yield of CD34+ cells. Protein microarray analysis revealed that there were eleven differentially expressed proteins in the NS group compared with the AA rat group. The identified specific NS might be implicated in rehabilitation of BMSC in AA rats, suggesting their potential of nutritional support in AA treatment.

Keywords: Aplastic anaemia: Nutritional supplements: Bone marrow-derived mesenchymal stromal cells: Rat models

Aplastic anaemia (AA) is a rare and life-threatening heterogeneous bone marrow (BM) failure disorder characterised by peripheral pancytopenia and marrow hypoplasia(1). Although the pathogenesis of AA has been associated with chemicals, drugs, radiation, infection and immune diseases, the exact mechanisms responsible for BM failure in AA are unclear. Recently, the main suggested underlying mechanism in AA is a primary haematopoietic stem cell (HSC) deficiency (HSC defect due to an abnormal balance between HSC death and differentiation), an immune-mediated inhibition of haemopoiesis, a defective BM microenvironment (BMME) or a combination of these conditions(2). Because the depression of haemopoiesis in AA is generally attributed to failure of HSC, it is necessary to explore a new stem cell therapeutic measure.

Mesenchymal stromal cells (MSC), also called mesenchymal stem cells, are multipotent progenitors. They were first isolated from BM and characterised by the ability to differentiate into adipogenic, chondrogenic and osteogenic lineages(3–5). Importantly, BM-derived mesenchymal stromal cells (BMSC) are an essential component of the BM haematopoietic microenvironment. As the abnormal alteration of MSC was observed in patients with AA, various studies have investigated the association between MSC and the pathogenesis of AA. MSC from AA patients have the typical MSC immunophenotype, aberrant cell morphology(6–8), as well as proliferative defects, lower clonogenicity and increased cell apoptosis(9,10). Jiang et al.(11) also showed that the fibroblast growth factor 1 treatment induces the proliferation of BMSC in patients with AA.

Abbreviations: AA, aplastic anaemia; BM, bone marrow; BMD, bone mineral density; BMSC, bone marrow-derived mesenchymal stromal cells; Epo, erythropoietin; HSC, haematopoietic stem cells; IFN-γ, interferon γ; MSC, mesenchymal stromal cells; NGF, nerve growth factor; NS, nutritional supplement; RELM-β, resistin-like molecule β; SD, Sprague–Dawley; SN3, serum from AA rat group with treatment of high-dose NS.

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Moreover, the differentiation capacity of AA-derived BM MSC appeared defective\(^{6,9}\). Treatment with arsenic trioxide partially reversed the differentiation imbalance. Arsenic trioxide could inhibit adipogenic differentiation, but promote osteogenic differentiation in AA BMSC\(^{6,12}\). However, the underlying mechanism of the influence of BMSC on AA treatment by nutritional supplement (NS) remains unclear and requires further investigation.

Use of natural health products to prevent, inhibit and reverse diseases is gaining increasing importance. Scientific evidence showed that a nutrient mixture, containing ascorbic acid, lysine, proline and green tea extract inhibited matrix metalloproteinase expression and mediated cell invasion and apoptosis in human falcioni anaemia fibroblast cell lines\(^{13}\). We previously developed strategies to accelerate the rehabilitation of AA mice by promoting peripheral blood cells production, marrow nucleated cells proliferation and mitochondria formation, increasing primary haematopoietic progenitors as well as repairing liver and spleen damage of AA mice\(^{14,15}\). In the present study, we examine the effect of NS (containing various kinds of amino acids, nucleotides, vitamins and metal elements) on BMSC from AA rats, evaluating proliferative behaviour of BMSC, suppressing BMSC differentiation and analysing protein expression.

**Methods**

**Nutrient mixture**

The nutrient mixture was supplied by Dalian Golden Axe Biological Technology Development Co., Ltd and made of lysine (6.6%), methionine (8.8%), phenylalanine (7.7%), threonine (3.3%), tryptophan (2.2%), arginine (8.38%), histidine (5.5%), glycine (3.3%), aspartic acid (3.3%), leucine (3.3%), isoleucine (3.3%), valine (4.4%), serine (2.2%), glutamine (3.3%), taurine (2.2%), whey acid (3.3%), nucleotide (4.4%), niacin (0.13%), folate (0.0099%), inositol (0.066%), Soyabean phospholipid (13.25%), vitamin A (0.0066%), vitamin C (1.1%), vitamin D (0.000066%), vitamin B\(_2\) (0.022%), vitamin B\(_3\) (0.022%), vitamin B\(_12\) (0.000044%), Fe (0.11%), Zn (0.11%), Mn (0.088%), Cu (0.022%), Se (0.001%), Cr (0.0002%), K (3.3%), Ca (3.3%), Mg (2.2%) (online Supplementary Table S1). According to the proto-prescription, oral application was prepared.

**Animals and treatments**

In order to induce an AA model, 6-week-old healthy male Sprague–Dawley (SD) rats (n = 126; mean body weight 191.1 ± 16.2 g) were obtained from Animal Facility of Dalian Medical University. All of experimental procedures were approved by the Animal Studies Ethics Committee of the Dalian Medical University, China. Rats were housed in animal facility with temperature and humidity setting to 24 ± 1°C, 55 ± 10% and with a 12 h light–12 h dark cycle. In all, 126 SD rats were divided randomly into five groups: control group, AA model group and three AA groups with different dose of the nutrient mixture supplement. The 1st day, AA rat models and the three different doses of the nutrient mixture groups were irradiated by X-ray (2.5 Gy). Then the rats were treated with intraperitoneal injections of cyclophosphamide (35 mg/kg, Sigma) and chloramphenicol (35 mg/kg, Sigma) in the 4th, 6th, 8th days. These steps were repeated the 15th day. The control animals were treated with physiological saline. After the 5th day, the rats of nutrient mixture groups were fed with nutrient mixture supplement in a dose-dependent manner (2266 ± 95, 1511 ± 3, 1057 ± 91 mg/kg d, according to Chinese Medical Dictionary) by intragastric administration. All rats received a standard diet during the experiment. After treatment with NS or physiological saline for 8 weeks, the rats were anaesthetised by isoflurane inhalation and decapitated for following experiments.

**Peripheral blood and bone marrow analysis**

The blood samples were collected from the SD rat eye sockets. A 0.5-ml aliquot of blood was anti-coagulated with 1.5 mg/ml dipotassium EDTA (Sigma). Blood samples were analysed with the Sysmex KX-21N Blood Cell analyser (Sysmex Corporation). Serum was isolated using serum separator tube (Microtainer; Becton Dickinson and Co.). The supernatants were collected for erythropoietin (Epo) and co-culture analysis. Epo levels were determined by an ELISA kit (no. ABIN566328; R&D Systems). The femoral marrow of SD rats was smeared. The femoral marrow cell suspension in 90% Ham’s F-12K (F-12K; Gibco) and 10% FBS (Gibco) was used to obtain the total BM nucleated cell count (femoral nucleated cell count (FNCC)) using the basophil channel of the Sysmex KX-21N.

**Isolation of bone marrow-derived mesenchymal stromal cells and haemopoietic stem cells**

BMSC and HSC were isolated from SD rats of each group. Rats were gently flushed out of pubis and femurs using F-12K and 10% FBS. Cells obtained from two pubis and two femurs were plated in 100 mm culture dish (BD Falcon) in 10 ml of BMSC media at 37°C with 5% CO\(_2\). After 24 h, non-adherent cells were collected for purification of CD34\(^+\) HSC. The remaining adherent cells, which were abundant in MSC, were trypsinised and replated until they achieved >85% density. Non-adherent cells were incubated with CD34-conjugated magnetic microbeads (Miltenyi Biotec) according to the manufacturer’s instructions. Afterwards, CD34\(^+\) populations were isolated using a separation column placed in a magnetic field, and were then subjected to phenotypic and functional assays with more than 90% survival rate determined by trypan blue staining.

**Phenotypic analysis by flow cytometry**

Surface markers (CD34, CD44, CD45 and CD90) of BMSC were identified using flow cytometric analysis. Approximately 1 × 10\(^5\) cells (after passage 3) were incubated in the dark for 30 min at 4°C with the fluorescein isothiocyanate (FITC)-conjugated antibodies against CD34 (sc-18917 FITC; Santa Cruz Biotechnology), CD44 (sc-18849 FITC; Santa Cruz Biotechnology), CD45 (sc-558477; BD Pharmingen), CD90 (sc-554894; BD Pharmingen), respectively. After washing with PBS, labelled cells were detected by using a flow cytometer (Beckman Coulter).
Co-culture assay

To explore the regulatory role of NS treatment in expansion of HSC, HSC were washed and resuspended in Iscove Modified Dulbecco Medium (Gibco), 50 ng/ml fetal liver tyrosine kinase-3 ligand, 50 ng/ml stem cell factor (SCF), 50 ng/ml thrombopoietin and 20 ng/ml IL-3 (all from PeproTech). HSC were then prepared into suspensions at a concentration of 1 × 10^5/ml. Four culture systems comprised this assay: (1) HSC suspensions plus 20% serum from the healthy control group (HSC + SC), (2) HSC plus 20% serum from the AA rat group with treatment of high-dose NS (HSC + SNS), (3) HSC plus MSC and SC. HSC suspensions were seeded on a confluent MSC layer with 20% SC, (4) HSC plus SNS and MSC. HSC suspensions were placed on a confluent MSC layer with 20% SNS. The cells were incubated at 37°C in 5% CO₂ for 7 d. Subsequently the expansion of HSC was analysed by flow cytometry.

Cell proliferation assay

The suspension of BMSC (1 × 10^5/well) were plated in ninety-six-well plates at a concentration of 100 µl of F-12K medium containing 10% FBS. The cells were then cultured by incubation at 37°C in 5% CO₂ for 2, 4, 6, 8, 10, 12 and 14 d. 10 µl CCK-8 (Sigma-Aldrich) solutions were added to the well for 2 h. The absorbance at 450 nm was immediately measured using a microplate reader. The results were recorded for further comparison.

Colonies formation assay

To measure focus formation, cells (1 × 10^5) were trypsinised to single-cell suspension and seeded in six-well plates. The cultures were maintained in the F-12K containing 10% FBS, with medium changes every 3 d. At day 14, the colonies were stained with 0.2% crystal violet, and foci were counted. Images of the colonies were obtained using a NIKON digital camera.

Osteoblast differentiation and alizarin red staining

Osteoblast differentiation was performed in cells (2 × 10^5/well) plated in twelve-well plates in osteoblast-induction media containing Dulbecco’s modified Eagle medium (DMEM) supplemented with 10% FBS, 10 µM β-glycerophosphate, 50 µg/ml L-ascorbic acid-2-phosphate and 10 nM dexamethasone (Sigma-Aldrich). The medium was changed every 3 d during induction period. Ca deposition, at day 14 of osteoblastic differentiation, was determined by BCA Protein Assay Kit (Pierce). Equal amounts of the protein (50 µg) were resolved by running gel (10% SDS-PAGE) and blotted onto same size nitrocellulose membranes. After being blocked for 2 h at room temperature in PBS containing 0.1% Tween 20 (PBST) and 5% skimmed milk, the membranes were hybridised with the following primary antibodies at room temperature for 2 h. Then, slides were washed twice with wash solution and incubated with diluted biotin-conjugated antibodies at room temperature for 2 h. After washing, slides were incubated with streptavidin-conjugated antibodies at room temperature for 2 h. Then, slides were washed and then reacted with streptavidin-conjugated fluor (104/ml) for 10 min. Images were captured using a digital camera system (Olympus DP12-2) coupled to an inverted optical microscope (Olympus CKX41). The number of adipocytes was counted and adipocyte content was calculated by summing up the number of adipocytes from three representative images.

Bone mineral density analysis

The bone mineral density (BMD) of the rats was determined using a dual energy X-ray absorptiometry system (Lunar iDXA; GE Healthcare) at the health check centre of the Second Affiliated Hospital of Dalian Medical University. The femurs were scanned at mid-point, and BMD was calculated according to the following formula: BMD (g/cm²) = the bone mineral content (g/cm) / bone width (cm).

Real-time PCR analysis

RNA was extracted using the RNeasy Mini Kit (Qiagen). The first strand complementary DNA was synthesised from 1 µg of total RNA using QuantiTect Reverse Transcription Kit (Qiagen). The primers were summarised in the online Supplementary Table S2. Real-time quantitative PCR was performed using SYBR-Green-quantitative real-time PCR Master Mix kit (Toyobo Co.). The data were normalised to the geometric means of the reference gene glyceraldehyde 3-phosphate dehydrogenase (GAPDH). The relative expression levels of each target gene were determined by using Biosystems 7300 Real-Time PCR system (ABI).

Protein microarray

Total protein was extracted from MSC. The protein concentration was determined by BCA Protein Assay Kit (KC-430; KangChen). Before processing, protein microarray slides specific for rats (ninety proteins, CODE: PAM-G2; RayBiotech) (online Supplementary Table S3) were blocked with blocking buffer for 30 min and then incubated with samples for 1–2 h at room temperature. After washing, slides were incubated with diluted biotin-conjugated antibodies at room temperature for 2 h. Then, slides were washed and then reacted with streptavidin-conjugated fluor at room temperature. At last, slides were scanned by Axon scanner. By comparing the signal intensities, relative expression levels of cytokines were made. The intensities of signals were quantified by densitometry. Raw intensities were revised by background and normalised by median. Fold changes in protein expression were calculated.

Western blot analysis

The results from protein microarray were confirmed by Western blot analysis. Cells were lysed in RIPA buffer (Abcam). Total protein quantification was performed using BCA Protein Assay kit (Pierce) according to the manufacturer’s instruction. Equal amounts of the protein (50 µg) were resolved by running gel (10% SDS-PAGE) and blotted onto same size nitrocellulose membranes. After being blocked for 2 h at room temperature in PBS containing 0.1% Tween 20 (PBST) and 5% skimmed milk, the membranes were hybridised with the following primary
Nutritional supplementation and stromal cells

Statistical analysis

Each experiment was performed at least in triplicate, and the measurements were performed in three independent experiments. Data are expressed as means and standard deviations. The differences between the control and AA groups were evaluated by Student’s t test. Comparison of each NS-treated group with the AA group was determined by two-way ANOVA. $P < 0.05$ was considered statistically significant. All analyses were performed using SPSS 17.0 statistical packages (SPSS Inc.).

Results

Nutritional supplement effects on peripheral blood of aplastic anaemia rats

Peripheral blood parameters were first performed automatically in a haematology analyser. Leucocytes, erythrocytes, Hb and the mean platelet (Plt) counts were notably lower in the AA rat group than in the control group ($P < 0.05$ for all parameters, Table 1). In addition, the mean cell Epo of the AA group was significantly increased to 25-62-fold $v.$ that of the control group ($P < 0.05$).

After 8 weeks of treatment, peripheral blood parameters of rats from each group were determined. Leucocytes, erythrocytes, Hb and Plt counts were all increased significantly in the three NS groups compared with the AA rat group, respectively (Table 1). In addition to these increases, there was a significant reduction in Epo at this time ($P < 0.05 \ v \ AA$ rats). This result showed that NS indeed has effect on peripheral blood parameters and Epo level of AA rats.

Table 1. Nutritional supplement affects parameters of peripheral blood of rats (Mean values and standard deviations)

<table>
<thead>
<tr>
<th>Groups</th>
<th>EPO (mU/ml)</th>
<th>Leucocytes ($\times 10^9/l$)</th>
<th>Erythrocytes ($\times 10^{12}/l$)</th>
<th>Hb (g/l)</th>
<th>Plt ($\times 10^9/l$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>26</td>
<td>50.62</td>
<td>3.17</td>
<td>8.49</td>
<td>2.70</td>
</tr>
<tr>
<td>AA</td>
<td>25</td>
<td>1296.86†</td>
<td>155.78</td>
<td>1.74†</td>
<td>0.81</td>
</tr>
<tr>
<td>High dose</td>
<td>25</td>
<td>386.05*</td>
<td>69.37</td>
<td>7.39*</td>
<td>1.31</td>
</tr>
<tr>
<td>Middle dose</td>
<td>26</td>
<td>473.49*</td>
<td>85.16</td>
<td>4.79*</td>
<td>0.98</td>
</tr>
<tr>
<td>Low dose</td>
<td>24</td>
<td>699.01*</td>
<td>128.58</td>
<td>2.08</td>
<td>0.62</td>
</tr>
<tr>
<td>Control</td>
<td>26</td>
<td>50.62</td>
<td>3.17</td>
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<td>0.62</td>
</tr>
</tbody>
</table>

EPO, erythropoietin; Plt, platelet; AA, aplastic anaemia.

Nutritional supplement influences bone marrow of aplastic anaemia rats

To measure the effect of NS on BM of AA rats, the femoral marrow was smeared after NS treatment. The severe damage of BM, characterised by lacking haematopoietic cell islands, was observed in the AA rats. Nutrient mixture-treated mice showed a progressive increase in BM cells after 8 weeks feeding of NS (Fig. 1(a)). In addition, the average number of FNCC isolated from 1 ml of BM aspirate varied between groups and was significantly lower in AA BM compared with normal BM samples (Fig. 1(a)). There was a significant difference in the number of FNCC isolated from BM aspirates of untreated and treated AA rats with NS. Nutrient mixture of high dose markedly increased all BM cell population of AA rats (Fig. 1(b)). The results also indicated that NS increased the population of BM cells in AA rats.

Nutritional supplement promotes the proliferation of bone marrow-derived mesenchymal stromal cells in aplastic anaemia rats

To determine the potential role of NS affecting proliferation of BMSC, AA rats were treated with NS followed by BMSC isolation from mononuclear cells derived from BM samples. Flow cytometry analysis showed that the BMSC were positive for the specific surface markers CD44 (93-9.0%) and CD90 (95-1.1%), whereas negative for the haematopoietic markers CD34 (0-56%) and CD45 (0-77%) (online Supplementary Fig. S1). According to the results of the CCK-8 assay in Fig. 2(a), BMSC of AA rats grew slowly when compared with the control group. Interestingly, BMSC from high-dose NS were in a latent period of growth between days 0 and 2, exhibited logarithmic growth between days 2 and 6, and reached a plateau after day 14. Compared with BMSC of AA rats, the proliferation rate of BMSC of NS groups was increased, respectively. A time-dependent increase in proliferation rate was observed in all experimental groups, indicating the ability to maintain proliferation during long-term culture.

Subsequently, we assessed the colony-forming capacity of BMSC in AA rats after treatment with NS. As shown in Fig. 2(b), the total number of colonies was significantly lower in BMSC of AA rats compared with the control group. To compare the effects on AA rats with treatment of NS, we conducted the same series of analyses based on BMSC-derived colony formation, demonstrating that colony formation was increased by NS treatment in AA rats (Fig. 2(b)). Therefore, it was inferred that NS might promote the proliferation of BMSC in AA rats.
MSC and SNS markedly improved the yield of CD34+ cells than with control culture. Interestingly, co-incubation of HSC with from the HSC + SNS and HSC + MSC + SC groups as compared other three culture systems, implicating that NS promoted the expression of BMSC during their induction into osteoblast differentiation. The mRNA levels of fatty acid-binding protein 4 (FABP4), PPARG, adiponectin and CCAAT/enhancer-binding protein α (C/EBPα) were significantly increased in the AA group compared with the control (Fig. 4(a)). When BMSC were differentiated into adipocytes, the levels of these genes were significantly diminished in AA rats treated with NS. These data suggested that decreased expression of FABP4, PPARG, adiponectin and C/EBPα by NS suppressed adipocyte differentiation. Online Supplementary Fig. S3A and S3B show the influence of NS administration on BMD and adipocyte content of AA rats. Compared with the control group, AA rats displayed a decreased BMD of femur, accompanied by an elevation of adipocyte accumulation in MSC culture. As expected, the reduction in BMD was significantly improved and adipogenesis was inhibited in the group exposed to NS at high-dose. BMD of rat groups treated by middle and low dose was only slightly higher than that of the AA group, whereas the adipocyte

**Nutritional supplement promotes the proliferation of haemopoietic stem cells by affecting mesenchymal stromal cells**

CD34+ cells were enumerated using flow cytometry analysis over 7 d of culture. As seen in online Supplementary Fig. S2, there was a significant increase in the amount of CD34+ cells from the HSC+SNS and HSC+MSC+SC groups as compared with control culture. Interestingly, co-incubation of HSC with MSC and SNS markedly improved the yield of CD34+ cells than other three culture systems, implicating that NS promoted the expansion of HSC possibly by affecting MSC.

**Nutritional supplement promotes the differentiation of bone marrow-derived mesenchymal stromal cells into osteoblasts in aplastic anaemia rats**

To address the potential of NS to induce osteoblasts, we investigated differentiation of BMSC into osteoblasts. Formation of osteoblasts was decreased in AA rats compared with the control group (Fig. 3(a)). Furthermore, number of osteoblasts was significantly increased in AA rat treatment with NS compared with the control group, indicating increased osteoblast differentiation in dose-dependent manner of nutrient mixture. We examined the transcription factors alkaline phosphatase (ALP) and osteocalcin (BGLAP) involved in the osteoblast differentiation at a time interval of 14 d. The ALP and BGLAP mRNA of BMSC was considerably decreased in the AA group compared with the control group (Fig. 3(b)). Furthermore, NS in the AA rats exerted dose-dependent stimulative effect on ALP and BGLAP expression of BMSC during their induction into osteoblast differentiation. These data supported the effect of NS on inducing osteoblast differentiation of BMSC.

**Nutritional supplement suppresses differentiation of bone marrow-derived mesenchymal stromal cells into adipocytes in aplastic anaemia rats**

To determine whether NS regulates adipocyte differentiation of BMSC in AA rats, BMSC were induced. We demonstrated that adipocytes of AA rats were significantly increased compared with the control group on day 14 (Fig. 4(a)). In contrast, NS significantly inhibited the commitment of BMSC into adipocyte lineage, suggesting impaired adipocyte differentiation in dose-dependent manner of nutrient mixture (Fig. 4(a)). Thereafter, we analysed the expression of key adipocyte-specific genes during adipocyte differentiation. The mRNA levels of fatty acid-binding protein 4 (FABP4), PPARG, adiponectin and CCAAT/enhancer-binding protein α (C/EBPα) were significantly increased in the AA group compared with the control group (Fig. 4(b)). When BMSC were differentiated into adipocytes, the levels of these genes were significantly diminished in AA rats treated with NS. These data suggested that decreased expression of FABP4, PPARG, adiponectin and C/EBPα by NS suppressed adipocyte differentiation.
content of AA rats lowered obviously in response to NS treatment at middle and low dose.

**Discussion**

In the present study, to help understand the possible involvement of NS in the reduced haemopoiesis observed in the BM of AA rats, the basic properties of BMSC from AA rats were compared with those derived from AA rats samples treated with NS. Several
differences were identified. Our observations might be explained, at least in part, by a potential nutritional support in AA treatment.

AA is a rare disease in which the normal generation of erythrocytes and various white cell lines in the BM is disrupted. In this work, we developed an animal model of AA, by irradiation with X-ray and treatment with intraperitoneal injections of cyclophosphamide and chloramphenicol in rats. In order to confirm that we were in fact in the presence of an animal model of this haematological disorder, we assessed the parameters. As compared with the control group, AA rats showed significant reduction in peripheral blood leucocytes, erythrocytes, Hb and plt, even more severe decrease in femoral nucleated cell isolated BM aspirate, and increase in serum Epo, suggesting that we achieved X-ray, cyclophosphamide and chloramphenicol-mediated AA rat model.

Although numerous studies have documented their association with AA, investigations involving the role of NS in AA treatment are considered to be limited. Indeed, Jaako et al. (17) reported that amino acid L-leucine treatment significantly improved the number of erythrocytes, Hb concentration, BM cellularity and alleviated stress haematopoiesis in Rps19-deﬁcient anaemia mice. Payne et al. (18) demonstrated in their study that L-leucine can activate the mTORC1 signalling pathway and act as useful therapeutic agent for Diamond–Blackfan anaemia.

**Fig. 3.** Effect of nutritional supplement (NS) on the differentiation of bone marrow-derived mesenchymal stromal cells (BMSC) into osteoblasts in aplastic anaemia (AA) rats. (a) Alizarine Red staining was performed after 14 d of NS treatment. (b) quantitative PCR analysis of osteoblastic gene expression in BMSC induced to osteoblast differentiation by NS treatment. The expression of each target gene relative to that of glyceraldehyde 3-phosphate dehydrogenase (GAPDH) was calculated (*P < 0.05). Values are means and standard deviations of three independent experiments. b: Control; AA; high dose; middle dose; low dose; ALP, alkaline phosphatase; BGLAP, osteocalcin.

**Fig. 4.** Effect of nutritional supplement (NS) on the differentiation of bone marrow-derived mesenchymal stromal cells (BMSC) into adipocytes in aplastic anaemia (AA) rats. (a) Oil Red O staining and the area of mature adipocytes were shown after 14 d of NS treatment. (b) The adipocyte-specific gene fatty acid-binding protein 4 (FABP4), PPARγ, adiponectin and CCAAT/enhancer-binding protein α (C/EBPα), were analysed by quantitative real-time PCR. The expression of each target gene relative to that of glyceraldehyde 3-phosphate dehydrogenase (GAPDH) was calculated (*P < 0.05). Values are means (n = 3) and standard deviations and representative of three independent BMSC lines. b: Control; AA; high dose; middle dose; low dose.
Fig. 5. Nutritional supplement (NS) mediates protein expression of bone marrow-derived mesenchymal stromal cells (BMSC) in aplastic anaemia (AA) rats. Protein microarray containing ninety expressions confirmed proteins were probed with BMSC, as described in the ‘Methods’ section. (a) The image of protein microarray was tested by AA and NS-treated groups. Of these ninety differentially tested by the control and AA groups. Significant increases in the seventeen proteins of BMSC recognised from AA groups to controls were indicated. MIP, macrophage inflammation protein; CNTF, ciliary neurotrophic factor. (b) The image of protein microarray was tested by AA and NS-treated groups. Of these ninety differentially reactive proteins, eleven were lowly expressed in BMSC when compared with NS-treated groups for AA groups. Values are means and standard deviations of three independent experiments. RELM-β, resistin-like molecule β; GFR-z2, glial cell line-derived neurotrophic factor receptor α2; IFN-γ, interferon γ.

(DBAY)\(^{19}\). Yu et al.\(^{19}\) showed that the appropriate vitamin D supplementation could partly correct the immune dysfunction by strengthening signal transduction through vitamin D receptor in patients with AA. DBA was also associated with higher serum levels of vitamin B12 and folate, which would provide an efficient method for ruling out these deficiencies in the investigation of chronic anaemia\(^{20}\). Given that a single or a complex of several micronutrients supplementation is helpful in the recuperation process of AA, we prepared NS according to a novel formula of nutrient mixture with thirty-six ingredients, including different amino acids, nucleotide, vitamin, folate, soyabean phospholipid, inositol, iron, zinc, manganese, etc., which was added in the daily diet of AA rats, subsequently the effect and underlying mechanism were explored. After 8 weeks of NS administration, AA rats showed a reduction in erythrocytes, leukocytes, PLT numbers and the Hb concentration, and these parameters were significantly improved on NS treatment. Whereas Epo was reduced in AA rats treated with NS. In addition, NS had effect on BM nucleated cell recovery in AA rats. The hypocellular blood and BM in AA rats were significantly improved on NS treatment, providing an excellent approach to test novel therapeutic agents enhancing AA rat haematopoiesis.
AA patients have a hypocellular BM which is replaced by abnormal BM, likely of mesenchymal origin, further supporting a potential contribution of the BM microenvironment to the pathogenesis of AA(40). It has been shown in several studies that BMSC played a vital role in modulating BM microenvironment and supporting haematopoiesis(21). Graft versus host disease is a complication commonly occurred in patients with haploidentical haematopoietic stem-cell transplantation. However, clinical practice revealed that co-transplantation of MSC can sustain haematopoietic engraftment and decrease the risk of graft failure(22). Moreover, MSC in AA patients exhibited significantly reduced haematopoiesis-supporting capacity(23). We, therefore, hypothesised that BMSC might contribute, directly or indirectly, to the pathogenesis of AA. In the present study, our data indicated that BMSC from AA rats had impaired functional properties in proliferation and colony-forming capacity. As expected, NS rescued the inhibitory effect of BMSC growth in AA rats, suggesting that proliferative behaviour of BMSC was mediated at least in part by NS treatment. Furthermore, a co-culture assay revealed that co-incubation of HSC with MSC and NS markedly improved the proliferation of CD34+ cells than the HSC + SNS or HSC + MSC groups, implicating that NS promoted the expansion of HSC possibly by affecting MSC.

In general, MSC possess the capacity to differentiate into various types of cells, repair tissue, modulate immune response and support haematopoiesis. MSC have been identified for their ability to differentiate into the bone, adipocytes and cartilage(24). And in some specific microenvironment, for example, injury, MSC could differentiate into the lung epithelial cells in lung injury(25,26). Therefore, efforts to identify the regulatory mechanisms that control the differentiation of BMSC into adipocytes or osteoblasts may lead to the development of new clinical applications and enhance our understanding of haematopoiesis, as BMSC are the primary sources of the haematopoietic microenvironment(27). Although NS play an important role in regulating proliferative behaviour of BMSC, its role in regulating BMSC differentiation of AA rats is unknown. In this study, research was initiated with the analysis of the association between NS and BMSC differentiation. The NS administration accelerated osteoblast, inhibited adipocyte differentiation and reduced accumulation of adipocytes in the AA group. Being a crucial component of haematopoietic niche, the osteoblastic niche has been known for its regulating role in HSC(27). However, few studies have focused on the effects of NS on BMSC differentiation and maintenance.

A co-culture system of MSC and differentiated osteoblasts could apparently improve the self-renewal capacity and ex vivo expansion of haematopoietic stem and progenitor cells (HSPC)(29). Conversely, marrow adipocytes inhibit the osteoblast differentiation of MSC by suppressing BMPs signalling(30). Our data gave the proof that NS was potential regulator of BMSC differentiation and therefore mediate haematopoiesis. In addition, NS exposure at high dose significantly enhanced BMD in AA rats, implicating the potential roles of NS in alleviating the damage of bone caused by AA.

As a major constituent of BM microenvironment, MSC can reconstitute the damaged stroma and secrete an array of haematopoietic cytokines, stem cell factor and extracellular matrix molecules to promote self-renew and differentiation(21,31). In this study, we used protein microarrays to profile protein expression in lysates obtained from BMSC. A total of ninety derivatives from MSC have been scanned for comparison and analysis. We compared protein profiles of BMSC in AA rats based on whether or not they had received NS treatment during the 8-week study period. However, MSC from the AA group with NS treatment secreted a diverse host of cytokines and haematopoietic growth factors at similar levels compared with those of the AA group, including differentiation and maturation-related molecules granulocyte colony-stimulating factor, macrophage-CSF, IL-1, IL-3, IL-6, as well as IL-12. The obvious dissimilarity of these factors was not observed between the AA group with or without NS administration. MSC from two groups also exhibited similar expression patterns of anti-inflammatory cytokines IL-10 and 13, adhesion molecules ICAM-1, HSPC homing promoter E-selectin and chemokines such as CCR4, CXCR4. Interestingly, of these ninety differentially reactive proteins, seventeen proteins were highly expressed in BMSC from the AA group when compared with the healthy control group, whereas eleven proteins had low levels in the AA group treated with NS compared with the AA group. Most of these altered factors are effective regulators for regenerative medicine. For instance, RELM-β belongs to a family of cysteine-rich secretory molecules initially described to control insulin resistance in rodents. RELM proteins have been shown to modulate inflammation and wound healing processes(32). Zanotti et al.(33) show that MSC are able to suppress activation of endothelial cells, angiogenesis and T-cell homing by releasing metalloprotease-1, thereby exerting anti-inflammatory effects. Other inflammation and immunoregulation-associated cytokines such as vascular endothelial growth factor (VEGF)(34), TGF-β(35) and its induced product, follistatin-like protein-1(36,37), were also changed following NS treatment in our study, which accounting for at least partially the regeneration-promoting potency of BM MSC induced by NS. In addition, MSC could be a source of regulators for signalling pathways and cellular interactions. In mouse skin, receptor associated late transducer (RALT) protein behaves as an inhibitor of epidermal growth factor receptor signalling(38). Two molecules, fractalkine (CX3CL1) and VEGF, have been shown to be involved in the process of homing and mobilisation of HSPCs(39). Interleukins play important roles in immune regulation and contribution to haematopoiesis. MSC themselves can produce therapeutic cytokines, such as IL-2(40). IL-2 has been reported to be increased in acquired AA(41), that was consistent with our findings. NGF plays an important role in the nervous system by supporting the survival and growth of neural cells, regulating cell growth, promoting differentiation into neuron, and neuron migration. β-NGF gene transfection promotes differentiation of BMSC into neurons through regulation of AKT and MAPK pathway(42). GFRα2 is described as a pituitary stem/progenitor cell marker involved in central and peripheral nervous system development(43,44). MSC from AA patients had deficient immune suppression of IFN-γ release(45). Increased production of IFN-γ has been confirmed to contribute to the pathogenesis of acquired AA, and immunosuppressive therapy may induce haematological remission by suppressing IFN-γ(46). We observed that above bioactive compounds released by MSC differed significantly between groups. With NS treatment at high dose, MSC also pronounced increased levels.
of HSC maintenance factors, including SCF and CXCL12. Thus, nutritional treatment may realise their ability to facilitate the recovery of BM of AA rats by regulating the production of cytokines from MSC.

In conclusion, our findings support the hypothesis that NS plays an important role in regulating the proliferation and differentiation potential of BMSC and contributes to haematopoietic supporting capacity in AA rats. Furthermore, in BM of AA rats, NS is involved in the protein expression of MHSC, which regulates the haematopoietic microenvironment. Identifying the regulatory mechanism of NS in BMSC may lead to the development of novel therapeutic approaches for AA.

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Supplementary material

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