Human Platelet Lysate Promotes Proliferation but Fails to Maintain Chondrogenic Markers of Chondrocytes

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ABSTRACT
Traditionally, foetal bovine serum (FBS) is used as a serum supplement for stem cell expansion in vitro. However, it is associated with xenoinmunisation and the transmission of animal pathogens, which may cause harm to stem cell recipients. As a safer alternative, human platelet lysate (HPL) has been introduced for propagating stem cells. Chondrocytes are expanded in vitro for cartilage repair via autologous chondrocyte implantation (ACI). In this study, we compare the efficacy of HPL prepared from expired platelet concentrates with that of FBS for promoting the proliferation and maintenance of the chondrogenic markers of primary human chondrocytes expanded in vitro. Chondrocytes were cultured in F12: Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 5% HPL, 10% HPL, and 10% FBS. The cell morphology, viability and growth rate were examined from passage 1 (P1) to P3. RNA was isolated from P3 cells for quantitative polymerase chain reaction (qPCR) to determine the gene expression level of the chondrogenic, dedifferentiation and hypertrophic markers. HPL promotes chondrocyte proliferation without compromising cell viability. In addition, the chondrocytes cultured with HPL were smaller. However, HPL failed to maintain the chondrogenic markers, except SOX 9, which was upregulated, but not significantly. Nonetheless, HPL also suppressed the expression of type X collagen (Col X), a chondrocyte hypertrophic marker. In summary, we demonstrate the benefits of HPL supplementation in human chondrocyte culture, where it enhances cell proliferation and suppresses chondrocyte hypertrophy. In the future, HPL can be used for the large-scale expansion of chondrocytes for ACI.

Keywords: Autologous chondrocyte implantation; cartilage; chondrocyte; osteoarthritis; platelet lysate

INTRODUCTION
Osteoarthritis (OA) is a common joint disease worldwide. The articular cartilage defects in OA are associated with pain and joint dysfunction, which severely affect patient quality of life (Hamoud et al. 2012; Ude et al. 2018). Apart from the elderly, OA is becoming more common in youths, especially those active in sports (Amoako & Pujalte 2014). Replacing arthritic cartilage with an artificial prosthesis has long been the treatment modality for severe OA (Ahmad et al. 2015). The usual treatment for older patients aged...
>60 years is total knee replacement. However, physicians have found it challenging to treat younger patients aged <50 years because the prostheses have a limited lifespan (Stiebel et al. 2014).

Autologous chondrocyte implantation (ACI) was introduced in 1987 as the first cell-based approach for treating articular cartilage defects (Viste et al. 2012). In brief, the technique involves isolating and culturing autologous chondrocytes from cartilage biopsy collected from the non-weight-bearing area of the joint and later implanting it in the damaged area under a periosteal cover. The periosteum provides a waterproof layer for covering the transplanted chondrocytes, as well as growth factors and mesenchymal stem cells that enhance cartilage regeneration (Peterson et al. 2010). ACI is an effective and durable choice for treating large, full-thickness cartilage defects of the knee joint, sustaining clinical and functional improvements for up to 20 years after the implantation (Peterson et al. 2010). Mistry et al. (2017) proved that ACI is not only cost-effective but also leads to long-term improvement. Scientists and clinicians have come to a consensus that ACI should be performed as early as possible and as the first surgical intervention to yield the best results (Sykes et al. 2018). All these findings suggest that ACI has the potential to be the definitive treatment for OA in the near future.

One of the crucial steps in ACI is the in vitro expansion of chondrocytes, which should be able to yield the needed cell number in the shortest time without compromising cell quality. However, dedifferentiation, which refers to the process whereby chondrocytes lose their chondrogenic markers and chondrocyte phenotype, is very common in cells expanded in vitro (Ma et al. 2013). Dedifferentiated chondrocytes lose their rounded morphology and have reduced secretion of cartilage extracellular matrix (ECM) such as type II collagen (Col II) and aggrecan (ACAN) (Schulze-Tanzil 2009). Hence, dedifferentiation reduces the ability of culture-expanded chondrocytes to regrow functional cartilage upon transplantation.

Foetal bovine serum (FBS) has long been used as a serum supplement for in vitro chondrocyte expansion. FBS is produced from blood drawn from bovine foetuses (Hemeda et al. 2014). It is widely used in cell culture because it is easy to produce, rich in paracrine factors that support cell proliferation and is suitable for a variety of cells (Yao & Asayama 2017). Despite its numerous advantages, its usage also carries a few disadvantages that render it not ideal for culturing cells meant for clinical use. The disadvantages of FBS include imprecisely defined content as well as the presence of animal pathogens and proteins that may cause immune activation in transplant recipients (Cimino et al. 2017).

In vitro cell expansion using human platelet lysate (HPL) was first introduced in the 1980s (Doucet et al. 2005). HPL can be produced from platelet concentrates through a simple freeze-thawing method. Similar to FBS, HPL is also rich in growth factors that support cell proliferation. The usage of HPL in cell culture lowers the risk of immune rejection and infection in transplant recipients. However, HPL presents a transmission risk of blood-borne diseases, as do other blood products used for transfusion. Typically, HPL is prepared from a large pool of platelet concentrates from multiple donors to minimize lot-to-lot variation (Hemeda et al. 2014).

The aim of the present study was to determine the growth rate and chondrogenic markers of primary human chondrocytes cultured with HPL prepared from expired platelet concentrates. The cell morphology, viability and growth rate were examined. In addition, the expression of chondrogenic, dedifferentiation and hypertrophic markers of cells cultured with the different growth supplements was quantified using quantitative polymerase chain reaction (qPCR).

MATERIALS AND METHODS

ETHICAL APPROVAL

The study was conducted with approval from Universiti Kebangsaan Malaysia Research Ethics Committee (Reference number: UKM PPI/111/8/JEP-2018-100).

PREPARATION OF HPL

Expired platelet concentrates were obtained from the Blood Bank Unit of Universiti Kebangsaan Malaysia Medical Centre (UKMMC) and stored at -80°C before processing. To prepare HPL, 3-5 bags of expired platelet concentrates were thawed in a 37°C water bath and pooled. The freeze-thaw cycle was repeated to ensure lysis of all platelets. Subsequently, the supernatant was collected and filtered through a 100-μm cell strainer (Corning, USA) after 15 min centrifugation at 4500 g. Heparin (final concentration, 4 IU/mL) was added to prevent clotting, and the processed HPL was stored at -20°C until used.

CHONDROCYTE CULTURE

Human chondrocytes were isolated from the cartilage of consenting donors who had undergone total knee replacement at UKMMC. The tissue was minced and treated with type II collagenase (Worthington, USA) to dislodge the chondrocytes. The chondrocytes were cultured in 6-well tissue culture plates in equal volumes of Ham’s F12 medium (Gibco, USA) and Dulbecco’s modified Eagle’s medium (Gibco) supplemented with 10% FBS (Sigma, USA), 200 mM L-glutamine (Gibco), 100 U/mL penicillin (Gibco), 100 μg/mL streptomycin (Gibco), 0.25 μg/mL amphotericin B (Gibco) and 50 mg/mL ascorbic acid (Sigma). The culture was maintained in a 5% CO₂ incubator at 37°C. The culture medium was changed three times a week.

CELL MORPHOLOGY, VIABILITY AND PROLIFERATION

The morphological features of the chondrocytes cultured with HPL and FBS were examined every 3 days by capturing
images under an inverted light microscope (Leica, Germany). Cultured chondrocytes were grown to 80% confluence before being released from the culture plate using 0.05% trypsin-EDTA (Sigma) to quantify the cells and determine their viability using a haemocytometer and trypan blue dye. Viability was determined using the following formula:

\[
\text{Viability} = \frac{\text{Total live cells}}{\text{Total live cells} + \text{Total dead cells}} \times 100\%
\]

The chondrocytes were expanded three times from passage 1 (P1) to P3 with similar cell seeding densities and culture conditions. The cell growth rate, i.e. population doubling time (PDT), was calculated using the following formula:

\[
PDT = \frac{\log 2}{\log N_2 - \log N_1}
\]

where \(t\) denotes the time in culture (hours); \(N_2\) denotes the cell number at the end of the passage; and \(N_1\) denotes the cell number seeded at the beginning of the passage.

TOTAL RNA EXTRACTION

Total RNA of the cultured chondrocytes was extracted using Direct-zol™ RNA MiniPrep (Zymo Research, USA) according to the manufacturer’s instructions. The extracted RNA yield and purity were confirmed using a spectrophotometer. Total RNA was stored at -80°C after extraction.

REAL-TIME PCR

Real-time PCR was performed to quantitatively analyse the gene expression of chondrogenic (Col II, ACAN, fibromodulin [Fmod], SOX 9), dedifferentiation (Col I) and hypertrophy (Col X) markers of P3 chondrocytes. Primer 3 software and the GenBank database were used to design the primers (Table 1). The extracted RNA was converted to complementary DNA (cDNA) using iScript™ Reverse Transcription (RT) Supermix (Bio-Rad, USA). Reactions were run using iTaq™ Universal SYBR® Green Supermix (Bio-Rad) and the Rotor-Gene Q RT-PCR cycler (Qiagen, Germany) for 40 cycles. This series of cycles was followed by melt curve analysis to examine the reaction specificity. The expression level of each targeted gene was normalized to that of glyceraldehyde-3-phosphate dehydrogenase (GAPDH), and the chondrocyte differentiation index (the ratio of Col II mRNA expression to that of Col I, Col II/Col I index) was calculated.

STATISTICAL ANALYSIS

All data are presented as the mean ± standard error of mean and analysed using GraphPad Prism 7 (GraphPad Software, USA) with \(n = 6\) for cell growth and \(n = 3\) for RT-PCR. Statistical significance was calculated using one-way analysis of variance (ANOVA) followed by Tukey’s multiple comparison test, where \(p < 0.05\) was considered significant.

RESULTS

MORPHOLOGY OF CULTURED CHONDROCYTES

The primary human chondrocytes cultured with HPL had poor cell attachment at P0. Thus, we cultured the cells with FBS at P0 and then with HPL starting from P1. The cells were cultured in medium supplemented with 5% HPL, 10% HPL and 10% FBS until confluent, and then subcultured and grown again to confluence until P3. The cells cultured with FBS and HPL displayed fibroblast-like morphology at all passages. However, the cells cultured with HPL were smaller, longer and narrower compared to that cultured with FBS (Figure 1). In addition, the cells cultured with HPL.

| Gene                                      | Primer sequence (5'-3') | Product size (bp) | Accession number |
|-------------------------------------------|-------------------------|------------------|-----------------|
| Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) | F - caatgaccccttcattgacc R - ttaaattgagggatgtcctg | 160              | NM_002046.5     |
| Type II collagen (Col II)                 | F - gggagttaaatcagacagacaa R - atcatacaccagctctcag | 175              | NM_001844.4     |
| Aggrecan (ACAN)                           | F - aacagtctgggacattggtctgg R - gttgaagctgaggtggtcg | 189              | NM_001135.3     |
| Fibromodulin (Fmod)                       | F - ctttaaagtcctgctctgc R - tcaatggtcatctcgctgc | 144              | NM_002023.4     |
| SOX 9                                     | F - taagactcaccagacacca R - taagccattgtcattgctctc | 194              | NM_00346.3      |
| Type I collagen (Col I)                   | F - gtctaaaggtgctcattggt R - accagttcgccttcatctc | 128              | NM_00088.3      |
| Type X collagen (Col X)                   | F - ccaagcactgcactagtgtgtctc R - taatgtctggtggtcatctc | 119              | NM_00493.3      |
clustered together and overlapped when reaching higher confluence at all passages.

**VIABILITY OF CULTURED CHONDROCYTES**

No significant differences were detected in the viability of cells cultured with the different supplements at P1 and P2, whereby all groups showed viability of >90%. At P3, the cell viability of the FBS group was slightly lower (88.02 ± 2.91%) and was significantly different compared to chondrocytes cultured with 10% HPL (Figure 2(a)).

**GROWTH RATE OF CULTURED CHONDROCYTES**

From P1 to P3, the viable cell yield when the cultures were 80% confluent was approximately 2.0-4.6-fold higher in the HPL groups compared the FBS group. There were significant differences between the 10% HPL group and 10% FBS group at all passages and between the 5% HPL group and 10% FBS group at P1. Even though the viable cell yield was approximately 1.7-fold and 1.8-fold higher in the 10% HPL group compared to the 5% HPL group at P2 and P3, respectively, the differences were not statistically significant (Figure 2(b)). At all passages, the HPL groups had significantly shorter PDT and significantly higher population doubling number compared to the FBS group. In addition, there was a significant difference in the number of population doubling between the 10% HPL and 5% HPL groups at P3 (Figure 2(c) and 2(d)).

**GENE EXPRESSION OF CULTURED CHONDROCYTES**

The expression of chondrogenic markers, except SOX 9, was lower in the HPL groups compared to the FBS group (Figure 3). Significant differences were found only for Col II and ACAN expression between the HPL groups and FBS group. No differences were detected for Fmod and SOX 9. The expression of Col I, a chondrocyte dedifferentiation marker, was significantly lower in the HPL groups compared to the FBS group (Figure 4). Furthermore, the expression of Col X, a chondrocyte hypertrophic marker, was also significantly lower in the HPL groups compared to the FBS group (Figure 5). None of the supplements could maintain the chondrogenic markers of the cultured chondrocytes, as the Col II/Col I index was very low for all groups (Figure 6).

**DISCUSSION**

ACI has been a targeted therapeutic approach for OA patients, as it has shown encouraging results for repairing full-thickness cartilage defects (Mistry et al. 2017; Peterson et al. 2010). ACI safety and efficacy are very much dependent on the functionality of the cultured chondrocytes. The biggest challenge in chondrocyte expansion is the loss of the chondrocyte phenotype during serial monolayer culture in a process known as dedifferentiation. Dedifferentiation alters chondrocyte morphology and metabolism (Hamada et al. 2013). HPL has
been regarded as a safer alternative to FBS for the expansion of cells targeted for therapeutic use. Thus, in the present study, we investigated the proliferation and chondrogenic markers of primary human chondrocytes cultured with HPL prepared from expired platelet concentrates.

Here, there were few differences in the morphology of chondrocytes cultured with HPL and FBS. With both supplements, the cells appeared flattened and elongated. Pereira et al. (2013) and Sykes et al. (2018) reported a similar observation. They found that the phenotype of monolayer chondrocytes cultured with HPL changed from polygonal to spindle-shaped. Nonetheless, we found that the chondrocytes cultured with HPL, especially with 10%, were slightly smaller, longer and narrower. Furthermore, the chondrocytes cultured with HPL tended to cluster together and overlap when reaching higher

FIGURE 2. Viability and growth rate of primary human chondrocytes cultured with FBS and HPL (a) The cell viability was above 90% except for the P3 cells cultured with 10% FBS, (b) HPL increased the viable cell yield at all passages, (c) HPL shortened the population doubling time at all passages, and (d) HPL increased the number of population doubling at all passages (n=6). *, p<0.05 between groups. *, p<0.05 compared to the 5% HPL and 10% HPL groups

FIGURE 3. Gene expressions of chondrogenic markers. The expression of Col II (a), Fmod (b) and ACAN (c) were lower for the 5% HPL and 10% HPL groups compared to the 10% FBS group. The HPL groups have higher expression of SOX 9 (d) compared to the FBS group (n = 3). *, p<0.05 compared to the 10% FBS group
confluence. These observations are likely to be the reason for the higher cell yield in the HPL groups compared to the FBS group at similar cell confluency from P1 to P3. Cultures supplemented with HPL could yield up to 4-fold more cells. Several researchers have reported the ability of HPL to promote the proliferation of cultured chondrocytes (Hildner et al. 2015; Sykes et al. 2018). Interestingly, the cell clustering and overlapping in culture did not affect the cell viability, which was >90%. In addition, we found that the growth-promoting capability of 5% HPL was reduced in response to higher cell passages. However, the chondrocyte growth rate was maintained in the 10% HPL group from P1 to P3. Thus, even though 5% HPL supplementation is effective for promoting chondrocyte proliferation at lower passages, 10% HPL supplementation might be needed for chondrocytes at higher passages.

Chondrocytes secrete a myriad of ECM proteins to maintain cartilage tissue. Thus, we performed qPCR to quantitate the gene expression of ECM proteins to examine the characteristics of the cultured chondrocytes. The expression of major ECM proteins of articular cartilage, i.e. Col II, ACAN and Fmod, were decreased in the HPL groups compared to the FBS group. These findings indicate that HPL failed to maintain the chondrocyte phenotype. In contrast, SOX 9 expression was higher in the HPL groups. Similarly, Hildner et al. (2015) also reported higher SOX 9 expression in adipose-derived mesenchymal stem cells cultured with HPL compared to that cultured with FBS. SOX 9 is a transcriptional factor that regulates chondrocyte proliferation, cartilage formation and the suppression of chondrocyte hypertrophic changes (Sykes et al. 2018). Dedifferentiated chondrocytes with higher SOX 9 expression are more responsive to chondrogenic signals and produce more cartilage ECM (Hardingham et al. 2006; Li et al. 2004; Tew et al. 2005). We postulate that the chondrocytes cultured with HPL maintained higher SOX 9 expression due to the presence of a higher concentration of growth factors in the supplement. In the future, the growth factor responsible should be identified. Our RT-PCR results support the findings of Pereira et al. (2013), who reported that chondrocytes cultured with FBS lost the potential to redifferentiate faster than that cultured with HPL (Pereira et al. 2013). Similarly, Hildner et al. (2013) showed that chondrocytes expanded with HPL have higher Col II and GAG protein production when the cells were cultured in 3D (Hildner et al. 2015). In contrast, Kaps et al. (2002) and Sykes et al. (2018) found that HPL-expanded chondrocytes cultured in 3D secreted less GAG protein. The discrepancy in the results from these studies and ours might be due to the differences in HPL, as every study uses HPL prepared using different methods.

Higher Col I expression has long been used as an indicator of chondrocyte dedifferentiation. Col I is a fibril-forming collagen found in abundance in many tissues, including the cornea, dermis and tendons, but not articular cartilage (Law et al 2017a, 2017b, 2016). Here, Col I expression was lower in the HPL groups compared to the FBS group. Nonetheless, the Col II/Col I index was very low for all groups indicating that both supplements failed to maintain the chondrogenic phenotype of cultured chondrocytes (n = 3).
X expression was lower in the HPL groups compared to the FBS group. Col X is increased in chondrocytes in serial monolayer culture (Lin et al. 2008). These findings suggest that HPL supplementation might be able to prevent hypertrophic changes of cultured chondrocytes.

The limitation of this study is that our in-house HPL did not perform well in supporting the attachment of freshly isolated P0 chondrocytes. We were unable to establish primary chondrocyte culture for cartilage samples, with lower cell yield upon isolation using HPL supplementation. Thus, we used FBS to culture the P0 chondrocytes and used HPL only from P1 onwards. We suggest that in the future, a suitable coating substrate should be used to enhance chondrocyte attachment when HPL supplementation is used at P0.

CONCLUSION

Even though it failed to maintain the chondrogenic markers, HPL appears to be a better alternative to FBS for culturing chondrocytes, as it allows more efficient expansion and prevents hypertrophic changes of the cells.

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