Isolation, Culture and Characterization of Cancer Stem Cells from Primary Osteosarcoma

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Abstract:

Background:
Osteosarcoma cancer stem cells (CSCs) are defined as a subpopulation of osteosarcoma cells, which have the ability of self-renewal, proliferation and differentiation. This study aimed to identify CSCs from human osteosarcoma in vitro.

Methods:
Osteosarcoma CSCs were isolated and cultured with sphere-forming assay technique on an ultra-low well attachment surface plate. After sarcosphere colonies were formed, we conducted reverse transcriptase-polymerase chain reaction (RT-PCR) to detect the expression of genes of embryonic stem cells such as \textit{NANOG}, Oct3/4, STAT3, and gene of MSC CD133. Immunofluorescence analysis (IFA) of alkaline phosphatase (ALP), osteocalcin, and CD 133 was also performed to see the expression of osteosarcoma CSC surface protein with immuno-enzymatic staining principle. We also performed alizarin red staining to evaluate calcification in osteosarcoma CSCs.

Results:
The culture sphere-of the osteosarcoma cells showed three dimension round shaped colonies (sarcospheres) in slightly hypoxic and serum free condition which was not attached to the substrate with tight density. RT-PCR demonstrated that sarcospheres expressed genes which encode \textit{NANOG}, Oct3/4 STAT3, but not for CD 133. IFA showed positive protein expression of ALP, osteocalcin and CD 133 which was moderate, strong, and weak positive respectively. Sarcospheres also had a positive reaction toward alizarin red staining.

Conclusion:
Osteosarcoma CSCs could be isolated from human osteosarcoma by sphere-forming assay technique and characterized by the expression of genes of embryonic stem cells, such as \textit{NANOG}, Oct3/4, STAT3 and IFA of ALP, osteocalcin, and CD 133.

Keywords: Osteosarcoma, Cancer stem cells, Sarcosphere, CD133, \textit{NANOG}, Alkaline phosphate.

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1. INTRODUCTION

Osteosarcoma is a primary malignant bone tumor, commonly found in children and adolescents, which represents almost 60% of all bone sarcomas. It is an osteoid-matrix-producing solid tumor and mostly develops in long bones, usually on metaphysis close to epiphyseal growth plate [1 - 3].

The incidence of osteosarcoma in all populations is around 4-5 per 1 million people. Incidence estimation of osteosarcoma increases to 8-11 per one million individuals per year at the age of 15-19 years. Osteosarcoma is known to have a bimodal age distribution with first peaked in young adults and the second in the elderly [2, 4 - 7]. Within a period of 13 years (1995-2007), there were 219 cases (16.8/year) at Cipto Mangunkusumo General Hospital, Jakarta with the largest distribution in the second decade [8].

The outcome for a patient with osteosarcoma has improved significantly from 25% to 60% as a result of administration of systemic chemotherapy and the development of surgical techniques [1 - 3]. The multi-disciplinary therapy has resulted in disease-free survival of 60-80% while allowing the use of functional limb salvage surgery in >80% of patients [4]. However, 20 to 40% of patients ultimately die of osteosarcoma because of pulmonary metastases [2, 9]. Unfortunately, micrometastases are often already present at the time of diagnosis [10].

The management of metastatic osteosarcoma is very difficult and current surgical intervention still does not guarantee life viability [11 - 14]. One of the major causes failure of the cancer cells to respond to the chemotherapeutic agents administered is drug resistance [9]. Chemotherapeutic agents often fail to completely eliminate cancer stem cells (CSCs), which are able to repopulate the tumor mass, causing relapse and distant metastases. Thus, CSCs represent a pivotal pharmacological target to obtain effective therapeutic responses in cancer [15, 16].

Osteosarcomas are characterized by intratumor cell heterogeneity that could be explained by the recent CSC hypothesis. According to this theory, a subpopulation of cancer cells with stem-like properties, therefore named CSCs, is present within the hierarchical tumor organization. CSCs can divide asymmetrically, producing an identical daughter cell and have a pluripotent characteristic with self-renewing and the ability to differentiate into various cell types with high survival rates [17 - 20]. CSCs are derived from stem cells or progenitor cells that underwent dysregulation in proliferation and differentiation [21]. They are responsible for tumor development and recurrence, and drug resistance to chemotherapeutics [22 - 24].

Gibbs et al., reported the existence of CSCs in osteosarcoma [24]. They demonstrated that human osteosarcoma had a small number population of cancer cells that might grow in a clonal cluster environment when placed in a solution of serum-free conditions. The cell cluster is now referred to as sarcosphere or osteosphere [24].

A number of cell lines have successfully been cultivated and characterized from human [25 - 28] and canine [29, 30] osteosarcomas. A large number of human osteosarcoma cell lines are commercially available, like HOS [31], U2 OS [32], Saos-2 [33], or MG-63 [34]. CSCs of human osteosarcomas, in Indonesia, have rarely been studied, and to our knowledge, this work presents the first establishment and description of a human osteosarcoma cell line from primary human osteosarcoma.

In this study, cancer stem-like cells have been isolated from human osteosarcoma primary cultures and biomolecular characterization of these cells has been performed such as *NANOG*, Oct3/4, and STAT3), and CD133. The cultivated osteosarcoma cells were also tested for bone tumor markers as such as tissue unspecific alkaline phosphatase, osteocalcin, and alizarin red staining.

2. MATERIALS AND METHODS

2.1. Isolation and Culture of Human Osteosarcoma Cells

Primary tissue samples were taken from osteosarcoma patients who underwent open surgical biopsy or amputation. Each tissue sample was divided into two parts, one part for cell culture and one part for histopathologic examination. Tissue samples that were not intended for histopathologic examination were placed in sterile tubes containing DMEM with 2% antibiotic (Dulbecco's Modified Eagle Medium; Gibco®, Life Technologies, InvitrogenTM, Carlsbad, CA, USA) and then sent to culture laboratory immediately.

We cut the surgical samples into 2 – 4 mm square sections and washed with sterile phosphate buffered saline (PBS, Life Technologies Inc., USA). Tumor cells were released from the samples by enzymatic treatment with 0.1%
collagenase (Sigma Aldrich, Irvine, UK). Samples were agitated 20 minutes. At the end of incubation, samples were centrifuged at 700xg for 10 minutes. The supernatants were discarded and 0.25% trypsin (Invitrogen USA) were added and incubated for 10 minutes to reach single cells.

Cells suspension were homogenized and filtrated using strainer 70 micrometer then 10 mL of DMEM and penicillin-streptomycin 1% (Gibco®, Life Technologies, InvitrogenTM, Carslbad, CA, USA) were added. The suspension was washed by centrifugation at 700 x g for 10 minutes. Cell population was counted with a hemocytometer.

Cells suspension was centrifuged again at 700 x g for 10 minutes. Growth medium DMEM with NaHCO₃ (Gibco®, Life Technologies, InvitrogenTM, Carslbad, CA, USA), 10% FBS and PS 1% was added. Cells were cultured in 37°C with 5% CO₂ and 95% air. The growth medium was replaced every 3-4 days. When the cells had reached 80% confluence, subculture was done by removing the cells from its substrate using 0.25% Trypsin - EDTA (Invitrogen,USA).

2.2. The Subculture of Osteosarcoma Cell

Cells were subcultured when the population reached ~80% confluence, the growth medium was removed, then monolayer cells adhered to the plate were rinsed with 5 mL PBS. 1 mL of 0.125% trypsin (Invitrogen,USA) was added to the plate and incubated at 37°C, 5% CO₂, and 95% air for 5 minutes. 1 mL growth medium was added to deactivate trypsin. The cells and medium were put into a 15 mL polypropylene tube and centrifuged at 400 x g for 5 minutes. The supernatant was removed, the cells were re-suspended with 10 mL DMEM. Subsequently, the cells were counted with a hemocytometer and were grown in TC flask 25 and incubated at 37°C, 5% CO₂ and 95% air.

2.3. Phenotypical Analysis: Morphological, Cytological and Immunocytochemistry Examination

Morphological examination of cells on a plate containing culture media was evaluated by using an inverted microscope and through cytology examination with standard Papanicolaou staining. Under light microscope Leica ICC50 HD®. Immunocytochemistry examination of cultured cells of alkaline phosphatase (ALP) and osteocalcin was conducted by immuno-enzymatic staining principle. The slide was rehydrated with graded alcohol (absolute, 96% and 80%) every 4 minutes, blocked with 0.5% H₂O₂ in methanol for 30 minutes and then washed in water for 5 minutes. Pretreatment of the slide was performed with citrate buffer in microwave Cook I and Cook II in 5 minutes, followed by blocking background target to block non-specific antigens and then incubated for 15 minutes. Then it was given primary antibody ALP / osteocalcin and incubated for 1 hour. The slide was given a universal ink secondary antibody to bind to the primary antibody for 15 minutes. After that, counterstaining was performed with haematoxylin for 1-2 minutes.

2.4. Isolation and Culture of Osteosarcoma CSCs: Sphere Forming Assay

The pleomorphic spindle cells (that were successfully isolated and cultured from primary osteosarcoma) were grown on 6 wells of ultra-low attachment surface tissue culture plate (Corning Inc, Corning, NY) with concentration of 1x10⁵ cells per well in DMEM that had been supplemented with FBS 10% and PS1% (Gibco®, Life Technologies, InvitrogenTM, Carslbad, CA, USA). Culture plate was covered with parafilm the first three days and incubated at 37°C, 5% CO₂ and 95% air. On day three, parafilm (3M, USA) was opened and the growth medium was replaced. Subsequently, the cells were put-in polypropylene 15 mL tube and underwent re-suspension with 4 mL medium. Centrifugation was performed at 500x g for 5 minutes (Beckman,Germany). The supernatant was removed, the cells were re-suspended with 1 mL medium and incubated at 37°C, 5% CO₂ and 95% air.

2.5. Sarcosphere Subculture

After the colonies were counted, the cells were put into a 15 mL polypropylene tube, centrifugation was conducted at 500x g for 5 minutes. The supernatant was removed and 3 mL 0.25% trypsin was added into the tube and incubated at 37°C, 5% CO₂ and 95% air for 5 minutes. Centrifugation was re-performed at 500x g for 5 minutes. Supernatant was removed, the cells were re-suspended with 2 mL medium. Cells were counted with a hemocytometer and were grown in ultra-low well attachment surface tissue culture plate with concentration 1x10⁴ cells/mL, covered with parafilm and incubated at 37°C, 5% CO₂ and 95% air.
2.6. Characterization of Osteosarcoma CSCs

Characterization of osteosarcoma CSCs was conducted with reverse transcriptase-polymerase chain reaction (RT-PCR) and immunofluorescence analysis (IFA). RT-PCR was used to detect the expression of pluripotent biomarkers such as *NANOG*, Oct3/4, STAT3 and mesenchymal stem cell such as CD133. IFA of ALP, osteocalcin, and CD 133 was also performed to see the expression of osteosarcoma CSC surface protein with immuno-enzymatic staining principle. Alizarin red staining was conducted to prove the calcification of sarcosperes.

2.7. Reverse Transcription Polymerase Chain Reaction (RT-PCR)

mRNA extraction using the RNeasy kit (Qiagen, USA) and RT-PCR using the Super Script® III Reverse Transcriptase kit (Invitrogen, USA) were done according to the standard procedure of the company. PCR amplification stages consist of denaturation at 94°C for 30 seconds, annealing (*NANOG*, Oct3/4, STAT3 and CD133 and GAPDH) at 54°C temperature for 30 seconds and elongation at 72°C for 30 seconds, repeated for 40 cycles. Sequence primer could be seen in Table 1. PCR results were passed into a 2% agarose gel that contained Ethidium Bromide in Tris-acetat-EDTA buffer solution (Gibco, USA). Electrophoresis was performed in 100 volts for 45 minutes and its visualization was documented by using Gel Doc (Biorad, USA).

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward Primer</th>
<th>Reverse Primer</th>
<th>Product (bp)</th>
</tr>
</thead>
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<tr>
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<td>GCTGAGATGCTCATCACAGGAG</td>
<td>ICGTTTCTTGACTGGGAGCGTTGC</td>
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</tr>
<tr>
<td>Oct ½</td>
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<tr>
<td>STAT3</td>
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</tr>
<tr>
<td>CD 133</td>
<td>AAGCATTGCGACATCTCTATGG</td>
<td>AAGCACAGAGGTCATTGAGA</td>
<td>233</td>
</tr>
</tbody>
</table>

2.8. Immunofluorescence Analysis (IFA)

Sarcospheres were were grown in 8-well tissue culture plate (CorningInc, Corning, NY). The cells were fixed in methanol for 2 minutes, then in acetone for an additional 2 minutes. Slides were rinsed three times using PBS (Invitrogen, USA). Following fixation, Primary antibodies for alkaline phosphatase (EPR4477) ab 108337 (Abcam, USA), osteocalcin (OC4-30) ab 13418 (Abcam, USA), CD 133/1 (MACS, Germany) were added to slides and incubated at 37°C temperature for 60 minutes. Slides were rinsed three times with PBS and secondary antibodies which had been labeled by FITC were added. Incubation was performed at 37°C temperature for 60 minutes. Evans Blue (Sigma, USA) staining was added to the slides and incubated at room temperature for 2 minutes. Slides were rinsed in PBS (Invitrogen, USA) viewed under a fluorescence microscope at 20x and 40x magnification (Nikon, Japan).

2.9. Alizarin Red Staining

Sarcospheres were cultured in tissue culture plate 12 wells (ComingInc, Corning, NY) for 7 days. After the growth medium was removed, each well was rinsed three times with PBS for 5 minutes. 10% of Formaldehyde was added to each well for 10 minutes, then the well was rinsed again three times with PBS. Alizarin red 2% was added into each well and incubated at 37°C for 15 minutes. Subsequently, each well was rinsed three times with PBS for 5 minutes and viewed with an inverted microscope (Nikon, Japan).

3. RESULTS

3.1. Osteosarcoma cell Cultures

Tissue samples were obtained from 2 osteosarcoma patients who had no neoadjuvant chemotherapy. Two primary human cell cultures were obtained from post-surgical tumor specimens with a histologically confirmed diagnosis of conventional osteosarcomas. Those cells showed a fibroblast-like morphology when grown as nucleated monolayer cells on plastic cell culture. On further observation, the cells underwent elongation with pleiomorphic shapes (Fig. 1).

The subculture of osteosarcoma cells was conducted after the cells reached 80% confluence. Passage P2 to P5 subculture was performed at day 7 but P1 waited for 10 days. We found that the total number of cells were increasing on each passage. This cell population has been successful to be subcultured up to the 5th passage, and may represent increased apoptosis, decreased proliferation since these cells were not immortal cells.
Fig. (1). Those showed the growth of cell colonies from isolated human osteosarcoma tissue. Spindle cells were attached to the culture plate surface (100x magnification). A). Cells on the culture media; B) and C). Some cells showed elongated appearance resemble fibroblast; D). The cells underwent elongation with pleiomorphic shapes; E). Those cells reached 80% confluent (x100).

Cytology examination with Papanicolaou staining showed osteoblast-like cells with pleomorphic, hyperchromatic nuclei and prominent nucleoli (Fig. 2). Immunocytochemistry examination demonstrated positive for alkaline phosphatase and osteocalcin expression (Fig. 2).

Fig. (2). Osteoblast like cells from osteosarcoma cell cultures with eccentrically located nuclei and prominent nucleoli. (A) (Papanicolaou, 400x). Immunocytochemistry examination demonstrated positive for alkaline phosphatase (B) and osteocalcin expression (C).

3.2. The culture of osteosarcoma CSCs: Sphere formation assay

We successfully isolated and cultured CSCs from human osteosarcoma cell cultures growing them in the stem cell serum free medium. These cells, representing a subset of human osteosarcoma cells possess stem cell-like features, retaining the ability to self-renew in vitro. Osteosarcoma stem-like cells showed three-dimensional sphere clusters (sarco-spheres) in slightly hypoxic condition in the first three days. Those osteosarcoma stem like-cells were not
attached (non-adherent) to substrate with tight density. Sarcospheres were able to produce normal adherent cell cultures (Fig. 3).

Fig. (3). Human osteosarcoma cells isolated and cultured in non-adherent culture/ultra low attachment surface tissue culture plate (100x magnification). A). On day (D)1 all spherical cells were not attached to substrate with tight density within the serum-free and hypoxic medium; B). On D7 The cells formed sarcospheres and reached ~ 80% confluence (P1); C). After sarcospheres transferred back into non-adherent culture plate, spherical cells were not attached to substrate again ; D) and E). Those cells reformed sarcospheres on D14 (P2) on D21 respectively.

Moreover, we observed that osteosarcoma stem-like cells, when transferred in the standard medium, are able to grow as adherent monolayers with the same characteristic of the corresponding osteosarcoma cell line of origin suggesting the retaining of ability to differentiate in non-stem cancer cells.

Moreover, we observed that osteosarcoma stem-like cells, when transferred in the standard medium, are able to grow as adherent monolayers with the same characteristic of the corresponding osteosarcoma cell line of origin suggesting the retaining of ability to differentiate in non-stem cancer cells.

A characteristic property of CSCs is their unique ability to self-renew. One of the methods to determine this ability is to test CSC capability of serial passage in vitro, thus to more deeply investigate the self-renewing ability of osteosarcoma stem-like cells, sarcospheres were dissociated into single cells and re-seeded at increasing cell densities, which allowed for the formation of separate non-adherent three-dimensional sphere clusters, in serum free medium. The passages were repeated every 7-21 days, by monitoring sphere formation efficiency. On day 7 sarcospheres reached ~80% confluence (Fig. 3A and B).
The osteosarcoma stem cells formed sarcospheres and were successful to be subcultured up to the 3rd passage. On passage 1, $5 \times 10^5$ cells were transferred back into ultra-low attachment surface tissue culture plates for each well with a total number of cells $2 \times 10^6$. On day 14, osteosarcoma stem-like cells formed sarcospheres with sphere formation efficiency being 0.030%. On day 21, passage 2, $2 \times 10^5$ those osteosarcoma stem-like cells formed sarcospheres with sphere formation efficiency as 0.03375%. On passage 3, those $9 \times 10^4$ cells were grown into ultra-low attachment surface tissue culture plates and formed sarcospheres as well. However, termination of the sarcosphere forming process was considered (Figs. 3C-E).

3.3. Characterization of osteosarcoma CSCs

3.3.1 RT-PCR examination showed that the sarcospheres expressed genes which encode pluripotent biomarker $NANOG$, Oct3/4, STAT3 and of MSC CD 133 (Figs. 4A and 4B). The RT-PCR result was supported by IFA examination. It demonstrated positive of ALP, osteocalcin and CD 133 which was moderate, strong, and weak positive respectively (Fig. 5). Calcification of the tumor cell cultures was determined by alizarin red staining. Sarcospheres also showed positive reaction toward alizarin red staining (Fig. 6).

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**Fig. (4A).** RT-PCR result of osteosarcoma CSCs. RT-PCR showed positive expression of pluripotent biomarkers $NANOG$, Oct3/4 and STAT 3. White band showed gene location targeted.

**Fig. (4B).** Osteosarcoma CSCs RT-PCR result using CD 133. RT-PCR demonstrated no expression of CD 133 marker.
Fig. (5). Immunofluorescence analysis of ALP, osteocalcin, and CD 133 was evaluated at 100x magnification. A). IFA of ALP showed moderate positive expression; B). IFA of CD133 demonstrated weak positive expression; and C). IFA of osteocalcin showed strong positive expression; (D) Negative control.

Fig. (6). We revealed that sarcospheres produced calcium in vitro. Alizarin red staining results on adherent culture plate. (A) On day 3; (B) On day 7. At 32x magnification.

4. DISCUSSION

Some literature stated that the culture of osteosarcoma cells, either from the primary cells as well as from a cell line, will show limited cancer cell population when grown in a serum-free on the non-adherent condition. Only primitive cells that have the ability to form sphere colonies that could survive [19, 35, 36]. Some researchers have managed to show that there are sarcoma cell group that have the capacity to form a sarcosphere in culture [6, 19, 35, 37].

A cell, which is regarded as a stem cell in a culture, should be able to demonstrate the capacity to proliferate, renew through a long period of time, and produce a large amount of derivatives that can differentiate into primary cells type [38]. A small amount of cells that have the stem cell ability will form a three-dimensional clonal patterns (spheres) in anoikis non-adherent cultured environment [16, 39]. The stem like cells can be expanded in vitro and repeatedly form spheres after multiple passages in long-term assays [16]. When those cells are transferred back into adherent culture plate, they may attach to the plate surface [35].

Those findings are consistent with our present study. In the ultra-low well attachment surface tissue culture plate, the cells had the ability to form sarcosphere. They could re-attach to the plate surface when the sarcospheres transferred back to a tissue culture plate. The ability to self-renew, proliferate by forming spheres and reattach to the monolayer culture plate, proved that the obtained cells contained small amount populations of primitive cells that were suspected as CSCs [6].
Also, similar results were reported by Gibbs et al., [19] who identified CSCs osteosarcoma from osteosarcoma tissue samples and cell line MG-63 through the sphere-forming assay. Wilson et al., [35] also reported CSCs from isolation and culture of canine osteosarcoma cell lines. Wang et al. [36] successfully observed human osteosarcoma cell lines. Gibbs et al., Wilson et al. and Wang et al. demonstrated a small amount of population of cells which were capable to form a sarcosphere.

Pluripotent cells are characterized by the expression of a group of genes required to maintain the stem cell-like feature such as \textit{NANOG} [40], Oct3/4 [41], and STAT3 [19], a transcription factor that regulates the suppression of differentiation genes [42], and although less evident CD133 as also described in canine osteosarcoma tissue biopsies [43]. Our study was consistent with those studies. RT-PCR demonstrated the positive expression of genes \textit{NANOG}, Oct3/4 and STAT3, but negative expression CD133. Also, a previous study by Gibbs et al., [19] reported the positive expression of \textit{NANOG} and Oct3/4 on sarcospheres. Wilson et al., [35] also successfully proved the expression of \textit{NANOG}, Oct3/4, and STAT3 in the adherent osteosarcoma culture and sarcosphere.Wang et al., [36] and Lou et al., [37] reported Oct3/4 expression in osteosarcoma and \textit{NANOG} expression in cell line MG-63 M respectively.

Osteosarcoma development may be ascribed to altered osteogenic differentiation into mature osteoblast [16]. In the present study, the sarcospheres demonstrated osteoblast activity. We revealed a subset of immunopositive cells which were consistent with the presence of markers of differentiation ALP and osteocalcin. It indicates an heterogeneous cell population not exclusively composed of completely mature cells but also by a more undifferentiated subpopulation.

A positive ALP also indicated the pluripotent trait was available on the sarcosphere. Singh et al., [44] and Pautke et al., [28] revealed positive expression of the ALP in human embryonic stem cells and in the osteosarcoma cell lines (MG-63, Saos-2 and U-2 OS) respectively. ALP activities were also found in canine osteosarcoma cell lines [27, 29,]. In human osteosarcoma cells, tissue unspecific ALP activity was dependent on cell density [28], therefore, this condition has to be considered when comparing data about ALP.

Osteocalcin, a small peptide hormone produced by osteoblasts, was used to verify the bony origin of our new cell lines. Osteocalcin was a protein marker of osteoblast that had been frequently used in MSCs studies [45, 46]. The use of osteocalcin marker in osteosarcoma cells was identical to the osteosarcomagenesis process [5].

Some previous studies have reported the various expression of CD 133 in osteosarcoma CSCs. CD133 is a glycoprotein with 3 isoforms that have been widely used for isolating CSCs. Di Fiore et al., [47] reported strong expression of CD133 in AB-OS cell line but not in osteosarcoma cell line MG-63. While Lou et al., [37] found a strong expression of CD133 osteosarcoma cell line MG-63-M but only slightly visible on osteosarcoma cell line MG-63. Saini et al., [48] found strong expression of CD133 in monolayer osteosarcoma cell lines (Saos-2, CHA59, Hu09) but weaker in the sphere Saos-2 and CHA59. A short study conducted by He et al., [49] mentioned that the positive expression of CD133 in osteosarcoma patient had clinical lung metastasis in 46 (65.7%) of patients. In this study, CD133 could not be detected by RT-PCR and showed a weak positive by IFA. We evaluated that the amount of osteosarcoma CSCs was too small. This could be demonstrated by FACS analysis before the cells formed sphere, CD 133 could be multiplied before analysis or with a qPCR technique.

We revealed that sarcospheres produced calcium \textit{in vitro}. Sarcosphere showed the bright orange-red color that indicated the occurrence of alizarin red complex bonded with calcium [41, 50]. Luk et al., [51] had reported the characterization of osteosarcoma cell lines Saos-2 and U-2 OS successfully. In the canine osteosarcoma, the sarcospheres also showed strong calcification after three days in culture, as demonstrated by alizarin red staining [50].

Osteoblast activities in osteosarcoma and osteogenic differentiated MSCs showed positive results on alizarin red staining [51]. Osteoblasts may be induced to produce extracellular calcium deposits in large numbers \textit{in vitro}. In this study, calcium deposits in sarcospheres showed bone formation \textit{in vitro} and could be stained with alizarin red. Alizarin red will bind with calcium to form alizarin red calcium complexes [52].

Although the methods utilized to detect the osteosarcoma CSC show populations with enriched stem cell-like characteristics, no specific markers for the osteosarcoma CSC have been established. Understanding of the MSC may aid in successful specific isolation of the osteosarcoma CSCs. Therefore, further research is required to fully understand the biological functions of CSC markers in order to accurately isolate osteosarcoma CSC [53]. The isolation and identification of the small subpopulation of CSCs from sarcomas was a long-term conundrum that stagnated research with no substantial breakthrough. Currently, the methods of isolations can be categorized into three main types, including sarcospheres (tumorospheres), side population sorting, and sorting cells according to specific cell surface
markers [54]. So far, the most useful and promising methods for selecting osteosarcoma stem cells are sarcospheres.

It is important to improve the methods of CSC isolation through the study of CSC biology, with the final goal of finding molecular targets and developing a very specific anticancer therapy directed against this particular cellular subset, which is probably responsible for the maintenance of the primary tumor, the development of its recurrence, and the origin of metastases in several organs. The study of CSC biology is also important to finding therapies that could be incisive in the cure of cancers, such as osteosarcoma, for which the survival rate after neo-adjuvant treatment remains very poor [53].

A host of different approaches are currently being explored to eliminate CSCs, ranging from small molecules that selectively inhibit key cell signalling pathways to immunological approaches, which can target multiple tumour antigens and potentially eliminating cancer cells through harnessing of innate immune responses.

CONCLUSION

Tissue stem cells have been identified through the ability of cells to form spheres and this method has also been used to assess CSC presence. CSCs could be isolated and cultured in vitro from human osteosarcoma with sphere forming assay technique. Sarcospheres showed three dimension round shaped colonies in slightly hypoxic and serum free condition which was not attached to the substrate with tight density. They expressed genes which encode of N/ANOG, Oct3/4, STAT 3 and had a positive protein expression of ALP, osteocalcin and CD 133, and also had a positive reaction toward alizarin red staining.

ETHICS APPROVAL AND CONSENT TO PARTICIPATE

This study was approved by the Health Research Ethic Committee Faculty of Medicine Universitas Indonesia- Cipto Mangunkusumo Hospital Reference no 752/UN2.F1/ETIK/2014.

HUMAN AND ANIMAL RIGHTS

No Animals were used in this research. All human research procedures followed were in accordance with the ethical standards of the committee responsible for human experimentation (institutional and national), and with the Helsinki Declaration of 1975, as revised in 2008.

CONSENT FOR PUBLICATION

Not applicable.

CONFLICT OF INTERESTS

The authors declare no conflict of interest, financial or otherwise.

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