Identification and characterization of metastasis-associated individualized gene expression signature in osteosarcoma

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Abstract
Introduction: Osteosarcoma (OS) patients with complete surgical resection still relapse with poor prognosis. Part of this is due to the inability to accurately detect distant metastasis. Thus it’s essential to identify metastasis-related biomarkers for OS.

Methods: In present study, a computational pipeline based on relative expression orderings (REOs) using gene expression profiles was constructed in metastases and non-metastases OS patients.

Results: 138 metastasis-associated gene pair signature (MGPS) were identified follow two independent datasets. In order to further extract metastasis-associated biomarker for clinical application, a metastases-specific co-expressed MGPS network was constructed and analyzed. MGPS such as MYL5 and RPL27A showed strong positive correlation (Cor =0.75, P <0.001) in metastatic OS patients. There were thirteen MGPSs in above network were associated with prognosis. These prognostic MGPSs could become as a specific classifier to distinguish metastases and non-metastases OS patients. Functional analysis showed MGPSs were associated with cancer metastasis-related functions. Drug and MGPS network could provide some drug candidates for treatment of OS.

Conclusions: Collectively, the roles of the MGPSs in OS were elucidated, which could be beneficial for understanding OS pathogenesis and treatment.

Background
The American Cancer Society reported that the incidence of childhood and adolescent cancers is increasing rapidly. Among these, osteosarcoma (OS) is one of the cancer with highest mortality rate in the paediatric population [1]. In the last 35 years, there was no significant improvement in the five-year survival rate if metastases are present during diagnosis [2, 3]. Similar to the observation in other cancers, metastasis is a major factor for most death (75%) of OS patients [4]. For non-metastatic OS patients, tumor resection together with chemotherapy is a successful treatment plan. However, planning appropriate treatment for metastatic OS is still a great challenge for researchers and clinicians.

Cancer metastasis include two process: (1) cancer cells disseminate from primary tumors to whole body. (2) establish secondary lesions in distant organs [5]. Identifying metastases-associated
biomarker was a key insight of diagnosis and treatment for OS. Recent years, some attempt has been contributed to identify metastases-associated biomarkers using gene expression profiles for many kinds of cancers [6-8]. However, the number of metastases-associated biomarkers which could be applicated for clinic is very small. Usually, most of the suggested metastases-associated biomarkers, classify patients into different risk groups by differential expression or clustering. The thresholds of differential expression which generated from training data set couldn't be directly applied to another data set. One of major reason is the gene expression level is sensitive based on diverse batch effect and platform of microarray. Furthermore, this means that the generated data of all the samples detected together should be normalized when only deal with a single sample. In other words, risk composition of other samples which are normalized together determines the risk prediction of a single sample [9].

One of insensitive approach for systematic biases of microarray is within-sample relative expression orderings (REOs). REOs are invariant and robust for monotonic data normalization and against differences within biological individuals [9, 10]. In past years, REOs have been proposed to identify prognostic signatures [11, 12]. However, the performance of REOs in predicting metastases-associated signatures for OS has been not reported. Therefore, it is worth trying to identify robust metastases-associated biomarkers for clinical application in OS based on the rank-based approach. In present study, a computational pipeline based on REOs using gene expression profiles was constructed in metastases and non-metastases OS patients. Follow two independent datasets, 138 common metastasis-associated gene pair signature (MGPS) were identified in OS. A metastases-specific co-expressed MGPS network and its topological characteristics was constructed and analyzed. Some of MGPS such as MYL5 and RPL27A showed strong positive correlation (Cor = 0.75, P < 0.001) in metastatic OS patients. Thirteen prognostic MGPSs in co-expressed MGPS network were discovered. These prognostic MGPSs could better classify metastases and non-metastases OS patients. Functional analyses suggested that MGPSs were related with regulation of cancer and cancer-related pathways. Drug and MGPS network could provide some drug candidates for treatment of OS. Collectively, this study could provide assistance for diagnosis and treatment of OS.
Results

**Extract GMPSs in OS for metastases and non-metastases patients**

We built an integrated and computational pipeline based on REO approach using gene expression profiles (Figure 1A). First, 245 metastases-related genes were identified and they would be used to follow analysis. Second, a 0-1 matrix for a pair of genes \( G_m \) and \( G_n \) in metastases and non-metastases OS patients according to expression pattern of the gene pair. Then, if the frequency of metastases OS patients accord with a particular REO pattern was measured by fisher’s exact test compared with non-metastases controls. The particular REO pattern was expression \( G_m > G_n \) or \( G_m < G_n \). Lastly, MGPSs were identified if the P values of Fisher’s exact test were smaller than 0.05. We performed this pipeline based on dataset from TARGET. The distribution of P values of MGPSs with fluctuation is between 0.01 and 0.05 (Figure 1B). We divide all P values into five regions: 0 to 0.01, 0.01 to 0.02, 0.02 to 0.03, 0.03 to 0.04 and 0.04 to 0.05. The amount in each region had no great difference (Figure 1C). In order to ensure the accuracy and stability of MGPSs for OS, another dataset GSE33382 from GEO was used for identifying MGPSs. The P values of GSE33382 showed similar distribution to TARGET dataset (Figure 1D, 1E). Lastly, we got 138 common MGPSs in both dataset TARGET and GSE33382 for OS (Figure 1F).

**Some MGPSs showed strong correlation in metastases-specific co-expressed network for OS**

The correlation values of MGPSs would showed great difference between metastases and non-metastases OS patients. According to the inference, a metastases-specific co-expressed network was constructed (Figure 2A). The network contained 71 nodes and 57 edges. More than half MGPSs showed great difference (absolute values of difference for PCCs > 0.25) between metastases and non-metastases OS patients. There were 50.88% and 5.26% MGPSs were both positive and negative correlated in metastases and non-metastases OS patients (Figure 2B). The correlated direction of 43.86% MGPSs were reversed in metastases and non-metastases OS patients. The global network showed scale-free distribution \( (R \text{ square}= 0.75) \) which is a specific topological feature of transcriptional regulatory network (Figure 2C). We found metastases-related gene RPL27A had
highest degree in this metastases-specific co-expressed network (degree= 19). Most of the interacted
strength between RPL27A and other genes had great difference in metastases and non-metastases
OS patients. For example, gene RPL27A and MYL5 had strong positive correlations (Cor= 0.75, P<
0.001) in metastatic OS patients (Figure 2D). However, gene RPL27A and MYL5 were not correlated
in non-metastatic OS patients (Cor= 0.02, P= 0.86, Figure 2E). The results indicated that these
MGPSs maybe play key and specific roles in metastases process for OS patients.

Some genes in MGPSs were associated with prognosis for OS patients

As we known, time to metastases and number of lesions are the most important prognostic factors for
OS patients [13]. Thus it’s essential to consider the prognostic roles of MGPSs in OS patients. 13
genes were associated with survival in above metastases-specific co-expressed network for OS
(Figure 3A). For example, prognosis-related MGPSs CYBB and P4HA1 showed strong negative
correlations in metastases OS patients (Cor= -0.35, Figure 3B). Prognosis-related MGPSs CYBB and
P4HA1 also showed strong negative correlations in metastases OS patients (Cor= -0.25, Figure 3C).
Gene P4HA1, CYBB and CD53 were all related with survival (Figure 3D, E, F). In addition, samples
with high expression for these genes usually showed worse survival. All the results indicated that
some genes in MGPSs were associated with prognosis for OS patients.

Prognosis-related genes in MGPSs could classify metastases and non-metastases OS
patients

We constructed a prognosis-related MGPS network which extracted from metastases-specific co-
expressed network. This network contained prognosis-related genes and their direct neighbored
genes in metastases-specific co-expressed network (Figure 4A). There were 25 nodes (13 prognosis-
related genes) and 18 edges in this prognosis-related MGPS network. We found three MGPSs including
P4HA1-CYBB, P4HA1-EVI2B and P4HA1-CD53 were all associated with survival. In order to validate if
these prognosis genes in MGPSs could become biomarkers for metastases of OS patients, we classify
metastases and non-metastases OS patients using gene expression profile follow a consensus
clustering method. The prognosis-related genes in MGPSs could distinguish all samples into diverse
groups. Final number of group was 4 based on area under CDF curve plot (Figure 4B and C). Each
OS patients group had a consensus expression pattern and could distinguish clearly (Figure 4D). Most OS patients could be classified accurately (Chi-square test, P< 0.001) and especially the last group (C4) were all matched with the non-metastases OS patients (Figure 4E). Collectively, all the above results indicated that the expression of prognosis-related genes in MGPSs could be served as specific biomarkers to distinguish metastases and non-metastases OS patients.

**Functional characterizations and drug repurposing candidates of MGPSs in OS**

Functional enrichment analyses were performed to characterize the functions of MGPSs in OS. For GO enrichment analyses, these MGPSs were enrichment in some critical biological functions such as regulation of microtubule polymerization, negative regulation of autophagy and negative regulation of smooth muscle cell migration (Figure 5A). Dynamics of actin filament cooperated with microtubules could drive cell motility process. Many studies revealed that microtubule dynamics were necessary for promoting epithelial-mesenchymal transition (EMT) [14-16]. Expansion of dormant tumor cells into metastases or anti-tumor inflammatory responses would be restricted and promoted by autophagy. On the contrary, metastasis would be promoted by self-eating based on strengthening fitness of tumor cell environmental stresses response including anoikis during metastatic progression [17]. For KEGG pathway enrichment analyses, these MGPSs were enrichment in some key pathways associated with cancer or metastasis such as Natural killer cell mediated cytotoxicity, B cell receptor signaling pathway and MAPK signaling pathway (Figure 5B). Natural killer cells participate to immune response against metastasis [18]. These functional enrichment analysis showed these MGPSs were associated with cancer metastasis.

We also constructed a drug-related MGPS network to explore drug repurposing candidates for OS (Figure 5C). We found Proline (DB00172) was a drug repurposing candidate and its target gene were P4HA1 and PYCR2. Proline is one of the twenty amino acids in organism, which is a component of protein. Normal functions of jonts and tendons and maintain and strengthen all reply on proline at a great extent [19]. Azacytidine is clinically used to treat myelodysplastic syndrome, a group of heterogeneous bone marrow stem cell diseases [20]. In our analysis, Azacitidine was considered as a drug candidate for OS. Azacitidine is a cytidine nucleoside analogue, which has the clinical activity of
myelodysplastic syndrome and the potential activity of solid tumor.

Discussion

In this study, we suggested that the MGPSs using REO approach follow expression level of genes play specific roles in metastasis process of OS patients. These MGPSs could be as metastasis-related biomarkers for OS. Differential expression analysis for gene expression profiles between metastases and non-metastases samples was considered as a common approach. The adoption of normalization for other samples contribute to differential expression of patients and it would generate a lot of uncertainty for risk classification of patient. Especially, the uncertainty would increase if the sample size was small and couldn’t represent all the disease patients [21]. Specially, qualitative REOs-based MGPSs would identify more accured personalized information of individual patient than traditional differential expression in clinical application.

A hypothesis that the OS patients with poor prognosis might harbor metastases was advanced. Thus we performed survival analysis and identified 13 prognosis-related genes in MGPSs. The problems of insufficient power and too complex to evaluate metastasis would be present based on all the MGPSs in OS patients. According the clinical needs, we proposed prognosis-related MGPSs based a strict voting criterion for survival analysis and proved that these prognosis-related MGPSs performed better than all the MGPSs. The results indicated that prognosis-related MGPSs maybe more suitable for clinical application of predicting metastasis. In order to explore if these prognosis-related genes in MGPSs could be specific biomarkers of metastasis in OS patients, we used them to classify metastases and non-metastases patients based on gene expression. The 13 prognosis-related genes could classify metastases and non-metastases patients (P < 0.001).

In prognosis-related GMPSs network, gene P4HA1 was a key node with highest degree. The previous study has reported that P4HA1 was the active catalytic component of prolyl 4-hydroxylase in cancers [22], glioma [23], prostate cancer [24] and pancreatic cancer [25]. P4HA1 can promote chemoresistance, tumor growth and metastasis. The prognosis-related gene CD53 could form a MGPS with P4HA1 in OS patients. CD53 is essential for CD2 signal transduction, growth regulation and cell survival of cancer [26]. P4HA1 was also a drug target in drug-related MGPS network. The roles of
P4HA1 in OS should be explored and validated.

Conclusions

In this study, some MGPSs were identified and characterized for OS patients based on gene expression profiles. Correlations of some MGPSs showed obvious difference between metastases and non-metastases samples. Prognosis-related genes in metastases-specific co-expressed network could become as specific biomarkers to classify metastases and non-metastases OS patients. The functional analysis showed the association between MGPSs and cancer metastasis in OS patients. Collectively, our study provides novel insights into the mechanisms underlying the roles of MGPSs in metastasis process for OS.

Materials And Method

**Clinical and gene expression profile datasets of OS**

There were two main independent datasets in present study. First dataset including gene expression and related clinical information of 92 OS patients containing 23 and 69 metastatic and non-metastatic patients were obtained from the Therapeutically Applicable Research To Generate Effective Treatments (TARGET, https://ocg.cancer.gov/programs/target) data portal, which included 17070 mRNAs. TARGET program provides a comprehensive genomic landscape to explore molecular characteristics of childhood cancers. Providing a novel guide for developing effective therapeutic plans based on generated data is the major goal of TARGET program. The case selection criteria and sample details could be obtained at https://ocg.cancer.gov/programs/target/projects/osteosarcoma. Then, the second gene expression profile was obtained from gene expression omnibus (GEO, https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE33382). Dataset GSE33382 gene expression profile included 23 and 69 metastatic and non-metastatic patients. The detailed information of patients could be found in previous study [27, 28]. Corresponding platform files (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GPL10295) was used to map the gene Ids to genes. The averaged expression values were calculated and applied when multiple probes corresponding to a same gene for each sample. The probes which couldn’t match any gene or matched not only a gene would be removed.
Identification of MGPS in OS

First, some metastases-related genes were obtained from CancerSEA (http://biocc.hrbmu.edu.cn/CancerSEA/), which is a database describes the functions of cancer cells [29]. We got 245 metastases-related genes and they would be used to follow analysis. The gene pair must have at least one metastases-related gene. Then, we built a 0-1 matrix for a pair of genes (G_m and G_n) in metastases and non-metastases OS patients. Lastly, if the frequency of metastases OS patients accord with a particular REO pattern was measured by fisher’s exact test compared with non-metastases controls. The particular REO pattern was expression G_m > G_n or G_m < G_n. The significant gene pairs evaluated with P < 0.05 were considered as MGPSs.

Construction and analysis of metastases-specific co-expressed MGPS network and its topological features

In order to construct a metastases-specific co-expressed MGPS network, pearson’s correlation coefficients (PCCs) are calculated for each MGPS in metastases and non-metastases patients, respectively. The MGPSs were considered as metastases-specific co-expressed MGPSs, if absolute values of difference between PCCs in metastases and non-metastases patients were bigger than 0.2. Then a metastases-specific co-expressed MGPS network was constructed using Cytoscape 3.3.0 (http://www.cytoscape.org/). The degree analysis of the network was also using Cytoscape 3.3.0.

Survival analysis of MGPSs for OS patients

In order to evaluate the performance about the MGPS for prognosis in OS, we performed survival analysis for genes in each MGPS. Follow the median value of expression level for each gene, the OS patients were divided into two risk groups. And then, Kaplan-Meier (K-M) survival analysis was used for the two groups. P < 0.05 was consider as prognostic gene for OS.

Classification power of the prognostic MGPS in OS

Consensus clustering approach was used to classify metastases and non-metastases OS patients based on expression data of genes [30]. A R package named ConsensusClusterPlus (https://www.r-project.org/) was performed to this process. Best category number was select when the areas of
Cumulative distribution function (CDF) curves were smallest. Chi-square test was applied to evaluate if metastases and non-metastases OS patients could be classified using this method (P<0.01).

**Functional and drug enrichment analysis for MGPSs in OS**

Online Enrichr (http://amp.pharm.mssm.edu/Enrichr/) tool was applied with default parameters to functional enrichment analysis for genes in MGPSs [31]. The enriched GO terms (P<0.01) and KEGG pathways (P<0.05) were extracted and considered as MGPS-associated functions. The gene-drug interaction data are download from DrugBank (https://www.drugbank.ca/) [32]. Then a drug-related MGPS network was constructed and analyzed to identify drug repurposing candidates for OS.

**Declarations**

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Not applicable.

**Disclosure of interest**

The authors declare that they have no competing interest.

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Not applicable.

**Availability of data and material**

The data that support the findings of this study are available from the corresponding author upon reasonable request.

**Ethics approval and consent to participate**

Not applicable.

**Patient consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

**Consent for publication**

Not applicable.

**Authors’ contributions**
ZGF conceived and designed the experiments, ZSR, LCB, FFG and LSH analysed the data, and ZYL, JCZ and ZBS wrote the manuscript.

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Figures
A. Extract significantly reversed gene pairs in OS for non-metastases and metastases samples

Gene expression profiles in OS for non-metastases and metastases samples

- 245 metastases-related genes

|       | M1 | M2 | ... | M22 | M23 | NM1 | NM2 | ... | NM22 | NM23 |
|-------|----|----|-----|-----|-----|-----|-----|-----|------|------|------|
| G1 > G2 | 0  | 1  | ... | 0   | 0   | 1   | 1   | ... | 0    | 0    |
| G1 > G3 | 0  | 0  | ... | 1   | 0   | 0   | 1   | ... | 0    | 1    |
| ......  | ...| ...|     | ... |     | ... |     |     | ...  |     |
| Gm > G1 | 1  | 0  | ... | 1   | 0   | 0   | 0   | ... | 0    | 0    |
| Gm > Gn | 0  | 1  | ... | 0   | 0   | 1   | 0   | ... | 1    | 1    |

Fisher's exact test

- P < 0.05
- Metastases-related reversed gene pairs

B. Density

C. P-values

D. Density

E. P-values

F. TARGET
   - 3848
   - 21553
   - 6788
   - 31773
   - 9309
   - 51218
   - 10522
   - 58581
   - 12498
   - 76791

GSE33382
   - 34
   - 0.01
   - 69
   - 0.03
   - 94
   - 0.02
Identification of MGPSs based on gene expression profile in OS patients. (A) The workflow of identifying MGPSs based on gene expression profile in metastases and non-metastases samples. (B) The density distribution curve shows distribution of P values of MGPSs based on TARGET dataset. (C) The box plots show P values of MGPSs based on TARGET dataset in diverse regions. (D) The density distribution curve shows distribution of P values of MGPSs based on GSE33382 dataset. (E) The box plots show P values of MGPSs based on GSE33382 dataset in diverse regions. (F) The venn plots show intersection between TARGET and GSE33382 datasets follow diverse regions of P values.
Construction of metastases-specific co-expressed MGPS network for OS. (A) A metastases-specific co-expressed MGPS network. Positive and negative correlations are represented by red and blue, respectively. The thicker edges represent bigger difference of correlations between metastases and non-metastases OS patients. (B) The pie chart shows percents for reverse, negative and positive interactions. (C) The plot shows degree distribution of metastases-specific co-expressed MGPS network. (D) The point plot shows the expression correlation of gene RPL27A and MYL5 in metastases samples. (E) The point plot shows the expression correlation of gene RPL27A and MYL5 in non-metastases samples.
Some MGPSs in metastases-specific co-expressed MGPS network were associated with survival in OS patients. (A) The radar chart shows the $P$ values of 13 prognosis-related MGPSs. (B) The point plot shows the expression correlation of gene P4HA1 and CYBB in metastases samples. (C) The point plot shows the expression correlation of gene P4HA1 and
CD53 in metastases samples. (D-F) The overall survival of two OS groups with high and low expression are showed in K-M curves. Two-sided log-rank test is used to evaluate the difference between two K-M curves. The risk score distribution of the genes based on gene expression. The patient survival status of the genes.
Prognosis-related genes in MGPSs could classify metastases and non-metastases OS patients. (A) Prognosis-related MGPS network for OS patients. The red circle represents prognosis gene. (B) Cumulative distribution function plot of the consensus index. (C) Relative change in area under CDF curve of different group number. (D) Consensus cluster heatmap of OS patients. (E) The gene expression heatmap, group type classified by consensus cluster is represented by sub label, the metastasis status of the OS patients is represented by sample type.
Figure 5

Functional characterizations and drug repurposing candidates of MGPSs in OS. (A) GO terms enriched for genes in MGPSs for OS and bar plots represent $-\log_{10}(P)$. The dot line graph shows number of enriched genes in each GO term. (B) KEGG pathway enriched for genes in MGPSs for OS and bar plots represent $-\log_{10}(P)$. The dot line graph shows number of enriched genes in each KEGG pathways. (C) The drug and MGPS network. Red and green represent drug and gene, respectively.