Supplementary manuscript of

Knowledge-Embedded Constrained Multiobjective Evolutionary Algorithm Based on Structural Network Control Principles for Personalized Drug Targets Recognition in Cancer

## Constructing PGIN by Paired-SSN method

For the paired-SSN method [S1], the first step is building the co-expression network based on the tumor sample and the normal sample of an individual patient [S2]. Then, we needed to determine whether this edge is used to construct the PGIN according to the P-value of the edge between gene *i* and gene *j* in the normal sample network and tumor sample network. The specific conditions are as follows: If the P-value is lower than 0.05 in the tumor sample network (the coexpression relationship between the interaction of two genes is significant) and larger than 0.05 in the normal sample network (not significant), or vice versa, this edge is retained to constitute the PGIN. In addition, we can get P-value of an edge by calculating  and then counting its Z-value of . The  of an edge between gene *i* and gene *j* and its Z-score can be calculated:



where *n* represents the number of reference samples and k represents the *k*-th patient in the perturbed network.  represents the PCC of an edge between genes *i* and *j* in the reference network; and represents the PCC of the edge between genes *i* and gene *j* in the perturbed network. Here, we calculated a measure to score the pPCC of edges in the PGIN by integrating gene mutation data across cancer type-specific data into the PGIN as follows:

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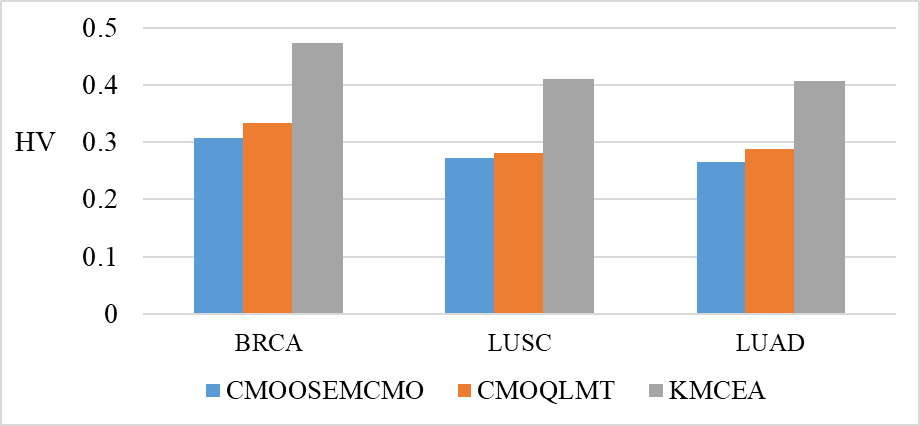
where norm represents the min-max normalized function.  and  respectively is the collection of tumors that exist mutated genes *i* and gene *j* after checking for somatic mutations in a given cancer data set;  indicates that 10% of the data falls under  after sorting a set of data in ascending order.

## The description of network and differential expression genes (DEG)-based methods

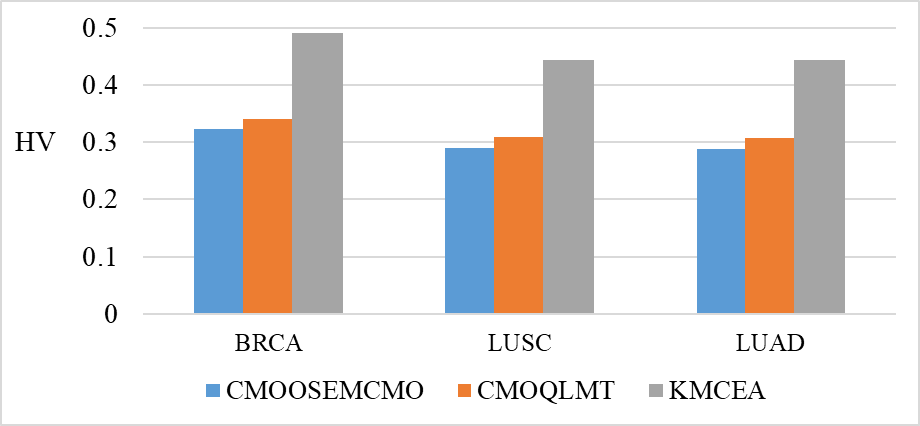
The network-based methods for identifying personalized drug targets (PDTs) were also taken for comparison in this paper, including CPGD [S2], ActiveDriver [S3], OncoDriveFM [S4], DriverML [S5] and Hub-genes. The cancer driver genes of CPGD and DriverMLwere obtained from their provided list of driver genes. Meanwhile, the driver genes in ActiveDriver and OncoDriveFM were obtained from the DriverDBv2 database [S6]. The hub gene selection method regards the hub genes in the constructed network as cancer driver genes. After the degree distribution of all genes T in the PGIN was obtained, a threshold was used in the following formula to obtain the hub genes:, where  and where  are the mean and standard variance of the degree distribution *T* of all genes, respectively. The DEG-based methods consist of DEG-Folchange, DEG-p-value, and DEF-FDR. Specifically, DEG-FoldChange selects the PDGs by calculating the fold-change between normal samples and tumor samples (log2(fold-change)| > 1). The DEG-p-value and DEG-FDR select the PDGs by calculating the p-value and FDR (<0.05) between a cancer tumor sample and a group of control samples, respectively.

## Compared results with two reinforcement learning-based CMOEAs

Two reinforcement learning-based CMOEAs are CMOOSEMCMO [S7] and CMOQLMT [S8]. CMOOSEMCMO uses deep reinforcement learning method to dynamically adjust the evolutionary operators of the population. CMOQLMT uses Q-learning and deep Q-learning methods to choose the most suitable auxiliary task during the evolutionary process. The compared results regarding HV and AUC results are provided in Figs. S-1 and S-2 respectively. Based on HV results, two compared algorithms are significantly worse than KMCEA on all three datasets under two frameworks. This is because two compared algorithms are designed for benchmark test functions. In addition, the AUC results also show that KMCEA is better than two compared algorithms. Therefore, the superiority of KMCEA is demonstrated.

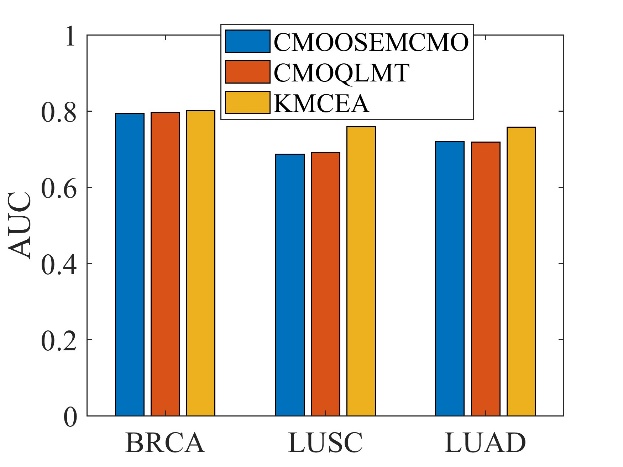
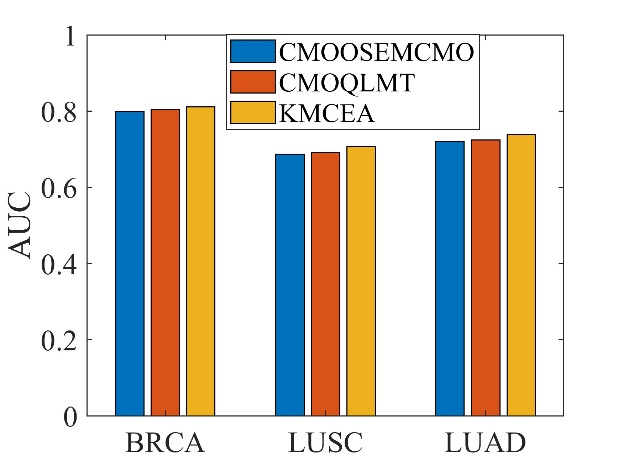


1. MMDS



1. MNCUA

Fig. S-1. Mean HV results of KMCEA and two compared algorithms on (a) MMDS and (b) MNCUA models.



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| --- | --- |
| 1. MMDS | (b )MNCUA |

Fig. S-2. Mean AUC results of KMCEA and two compared algorithms on (a) MMDS and (b) MNCUA models.

**References**

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