# An Adaptive Quantum-based Evolutionary Algorithm for Multiobjective Optimization 

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#### Abstract

An Adaptive Quantum-based Multi-criterion Evolutionary Algorithm called AQMEA is a new paradigm of decision making for complex systems. Quantum-based algorithms utilize a new representation for the smallest unit of information, called a $Q$-bit, for the probabilistic representation that is based on the concept of qubits. Evolutionary computing with Q-bit chromosomes has a better characteristic of population diversity than other representations, since it can represent linear superposition of states probabilistically. Moreover, we consider the three-criterion problem of task assignment.


Key-Words: - Quantum algorithms, multi-criterion optimization, distributed systems

## 1 Introduction

The foundations of infinite dimensional ordered spaces create the base for decision making among several criteria. They were statued at the turn of the century XIX and XX by Georg Cantor and Felix Hausdorff. In complex decision situation, some goals, criteria or players are in conflict. In fact, John von Neumann and Oskar Morgenstern noticed that an optimization problems in the context of a social exchange economy are "a mixture of several conflicting problems" that are "nowhere dealt with in classical mathematics" in their Theory of Games and Economic Behavior in 1944 [1].

Nowadays, there are several artificial intelligence techniques that can be applied to solve some multicriterion optimization problems [6]. Genetic algorithms, artificial neural networks, fuzzy logic algorithms, simulated annealing, tabu search, swarm approach and artificial immunological systems are crucial paradigms for a computer decision making [27].

On the other hand, a quantum-inspired algorithm is relatively new paradigm that can be applied for computer decision aid. Benioff considered a computer as a physical system and constructed a microscopic quantum mechanical Hamiltonian model of computer as represented by Turing machine [7]. It should be stressed that the Turing machine is a quite different approach for development of data processing than the von Neumann's paradigm. Feynman studied simulation models of physics for computer implementations what was the milestone to build quantum algorithms [13].

David Deutsch established the principles of quantum theory, and verified the Church-Turing principle to the universal quantum computer that is based on
a theoretical model named the quantum Turing machine [8]. Moreover, he introduced some quantum computational networks. Quantum computers accelerate the efficiency of calculations and do not allow computing functions that are not theoretically computable by classical computers due to the Church-Turing thesis: "Every function which would naturally be regarded as computable can be computed by the universal Turing machine" [9].

Shor proposed algorithms for quantum computation related to discrete logarithms and factoring [24]. In the factoring problem, it is given a composite number $N$ like $4,6,8,9,10,12$, and we want to find an integer $p$, strictly between 1 and $N$, that divides $N$. Shor's algorithm consists of a reduction of the factoring problem to the problem of order-finding what can be done on a classical computer. Then, a quantum algorithm is used to solve the order-finding problem. This approach is exponentially faster than the most efficient known classical factoring algorithm, and what is more, it can be applied to solve the widely used public-key cryptography method RSA that is based on the assumption that factoring large numbers is computationally infeasible for classical computers because no algorithm is known that can factor in polynomial time. However, Shor's algorithm shows that factoring could be efficient on a quantum computer. It was also a motivator for the development quantum computers and quantum algorithms.

In this paper, we consider a multi-criterion problem of task assignment, where a workload of a bottleneck computer and the cost of system are minimized. Furthermore, there are constraints for the performance of the distributed systems and the probability that all tasks meet their deadlines.

## 2 Quantum-based Algorithms

Vandersypen et. al., described an experimental implementation of Shor's quantum factoring algorithm using nuclear magnetic resonance [23]. They concluded that quantum computers, could factor integers in only polynomial time. Moreover, they reported an implementation of the simplest instance of Shor's algorithm: factorization of $N=15$ (whose prime factors are 3 and 5) by applying seven spin-1/2 nuclei in a molecule as quantum bits, which can be manipulated with room temperature liquid-state nuclear magnetic resonance techniques. This method of using nuclei to store quantum information is in principle scalable to systems containing many quantum bits. They demonstrated experimental and theoretical techniques for precise control and modeling of complex quantum computers and presented a simple, parameter-free, predictive model of de-coherence effects in system.

Hey described rules of quantum computing by development quantum phenomena, such a entanglement, superposition, and a quantum bit, to carry out operations on data. The basic principle is that these properties can be applied to represent data and execute instructions on these data [18].

Han and Kim introduced a genetic quantum algorithm and its application to combinatorial optimization problem [14]. They proposed a novel evolutionary algorithm, called a quantum-inspired evolutionary algorithm (QEA) characterized by a Q-bit as a probabilistic representation of the smallest unit of information. A Q-bit chromosome is defined by a string of Q-bits and it represents a linear superposition of binary states in search space probabilistically.

The Q-bit individual has higher population diversity than other known representations. A Q-gate is a variation operator that drives the chromosomes toward better solutions and toward a single state. Initially, there are various individuals represented probabilistically because a Q-bit chromosome represents the linear superposition of all possible states with the same probability. As the probability of each qubit converges either to 1 or 0 by the Q-gate, the Q-chromosome approaches to a single state and the diversity property vanishes progressively. An algorithm can treat the stability between exploration and exploitation.

Even though QEA is based on the idea of quantum computing, it is not a quantum algorithm, but a classical evolutionary algorithm. To demonstrate its numerical performance, experiments on the knapsack problem have been carried out. A quantum-inspired algorithm executed well devoid of premature convergence.

Then, Han, et. al., introduced a parallel quantuminspired genetic algorithm for combinatorial optimization problems [15]. A genetic algorithm operates on the population of chromosomes and that set
of Q-individuals can be divided on sub-populations performed by parallel set of quantum processors.

Han and Kim presented some results related to setting the parameters of quantum-inspired evolutionary algorithm for practical applications for a class of combinatorial optimization [16, 17]. That algorithm can be developed for the face verification [19] and for solving the Travelling Salesman Problem [26].

Recently, QMEA quantum-inspired multi-objective evolutionary algorithm has been proposed for multiobjective $0 / 1$ knapsack problems [21]. Experimental results pertaining to the multi-objective $0 / 1$ knapsack problem show that QMEA finds solutions close to the Pareto-optimal front while maintaining a better spread of non-dominated set. Another version of QMEA has been applied for a image segmentation [25]. Balicki proposed the other construction of QMEA for task assignment in computer networks [2].

In this paper, it is proposed an adaptive multiobjective evolutionary algorithm based on quantum computing, which is named an adaptive quantum-based multi-objective evolutionary algorithm (AQMEA). In the previous papers, quantum-inspired multi-objective evolutionary algorithm (QMEA) was proved to calculate better outcomes than conventional genetic algorithms.

To improve the quality of the non-dominated set as well as the diversity of population in multi-objective problems, an algorithm is proposed by employing the concept and principles of quantum computing such as uncertainty, superposition, and interference. We introduce the AQMEA to improve proximity to the Pareto-optimal front, preserving diversity intact by employing advantages of quantum-inspired evolutionary algorithm. The improving proximity means to find the better solutions which are evaluated as good individuals by fitness function.

In the multi-objective evolutionary algorithms are applied a strong elitist method with mechanisms to maintain diversity efficiently using non-dominated sorting and crowding distance assignment [22]. It is even more powerful if the elitism is further strengthened and the solutions are spread out by quantum mechanism. Multiple observations of Q-bit individuals allow a local search in the area of the non-dominated solutions.

Also, maintaining best Q-bit individuals in every generation can avoid the possibility of losing high quality individuals. Furthermore to deal with quantum computing concepts, the comparison mechanism is presented between the best group and the others. Convergence and preservation of diversity being the key issues under scrutiny, the proposed approach is expected to help improve the performance of AQMEA.

## 3 Rules of Adaptive Quantum-based Multiobjective Evolutionary Algorithm

An adaptive quantum-based multiobjective algorithm AQMEA utilizes a new representation, called a $Q$-bit, for the probabilistic representation that is based on the concept of qubits [20]. A qubit is a two-layer quantum system that can be modeled as the Hilbert space $H_{2}$ with the given base $B=\{|0\rangle,|1\rangle\}$. The Bloch sphere is a geometrical demonstration of the state space for a qubit and it may also refer to the space of an n-level quantum system.

A Hilbert space $H$ is a real or complex inner product space that is also a complete metric space with respect to the distance function induced by the inner product. It means that $H$ is a complex vector space on which there is an inner product associating a complex number to each pair of elements of $H$ that satisfies the properties of the complex conjugate, linearity, and positive definite. A real inner product space can be defined in the same way, apart from that $H$ is a real vector space and the inner product takes real values.

A qubit may be in the " 1 " binary state, in the " 0 " state, or in any superposition of the two [10]. The state $x_{m}$ of the $m t h$ qubit in the $Q$-chromosome can be represented, as follow:

$$
\begin{equation*}
Q_{m}=\alpha_{m}|0\rangle \oplus \beta_{m}|0\rangle, \tag{1}
\end{equation*}
$$

where
$\alpha_{m}$ and $\beta_{m}$-complex numbers that specify the probability amplitudes of the corresponding states,
$\oplus$ - a superposition operation,
$m$ - the number of the gene in the chromosome, $m=\overline{1, M}$.
Value $\left|\alpha_{m}\right|^{2}$ is interpreted as the probability that we observe the state " 0 ". Similarly, $\left|\beta_{m}\right|^{2}$ is the probability that state " 1 " is observed. A qubit may be characterized by the pair $\left(\alpha_{m}, \beta_{m}\right)$. There is, as below [11]:

$$
\begin{equation*}
\left|\alpha_{m}\right|^{2}+\left|\beta_{m}\right|^{2}=1 \tag{2}
\end{equation*}
$$

Let it be considered a four-bit register of a classical computer with 16 different four-bit strings 0000,0001 , $0010, \ldots, 1110,1111$. If it is a deterministic computer, then a four-bit register is in one of those states with probability 1.

However, if it is a probabilistic computer, then there is a possibility of it being in any one of a number of different states. We can describe this probabilistic state by sixteen probabilities $p_{0}, p_{1}, \ldots, p_{\mathrm{E}}, p_{\mathrm{F}}$. It means that a probabilistic computer is in one point from all possible states. There is a constraint that sum of these probabilities is equal to 1 .

The state of a four-qubit quantum computer is described by a sixteen-dimensional vector ( $\alpha_{0}, \alpha_{1}, \ldots$, $\alpha_{F}$ ), called a wave-function. However, the sum of the squares of the coefficient magnitudes, $\left|\alpha_{0}\right|^{2}+\left|\alpha_{1}\right|^{2}+$ $\ldots+\left|\alpha_{E}\right|^{2}$, must be equal to one. Moreover, the coefficients are complex numbers.

Since states are represented by complex wavefunctions, two states being added together will undergo interference. This is a main difference between quantum computing and probabilistic classical computing [10].

If we measure the four qubits, then we observe a four-bit string. The probability of measuring a string is equal the squared magnitude of that string's coefficients. Probability that we read state as 0000 is $\left|\alpha_{0}\right|^{2}$, probability that we read state as 0001 is $\left|\alpha_{1}\right|^{2}$, and probability that we read state as 1111 is $\left|\alpha_{F}\right|^{2}$. Thus a measurement of the quantum state with some complex coefficients ( $\alpha_{0}, \alpha_{1}, \ldots$, $\alpha_{F}$ ) gives the classical probability distribution $\left(\left|\alpha_{0}\right|^{2}\right.$, $\left.\left|\alpha_{1}\right|^{2}, \ldots,\left|\alpha_{F}\right|^{2}\right)$. We say that the quantum state "collapses" to a classical state.

In linear algebra, a basis plays important role and it is a set of vectors that, in a linear combination, can represent every vector in a given vector space, and such that no element of the set can be represented as a linear combination of the others. A basis is a linearly independent spanning tree. A sixteen-dimensional vector can be specified in many different ways, depending on a basis chosen for the space. The basis of four-bit strings is known as the computational basis, and it is suitable. However, the other bases of unit-length, orthogonal vectors can also be applied [11].

Bra-ket (Dirac) notation is used for describing quantum states by development of angle brackets and vertical bars. Moreover, it is less common used in mathematics because the inner product (or dot product) of two states can be denoted by a bracket, $\langle\alpha \mid \beta\rangle$ consisting of a left part, $\langle\alpha$, called the bra, and a right part, $|\beta\rangle$, called the ket [10].

Dirac notation is often used to make the choice of basis. For example, the state $\left(\alpha_{0}, \alpha_{1}, \ldots, \alpha_{F}\right)$ in the computational basis can be written as $\alpha_{0}|0000\rangle+\alpha_{1}|0001\rangle+\alpha_{2}|0010\rangle+\ldots+\alpha_{E}|1110\rangle+\alpha_{F}|1111\rangle$. We apply the notation $|0001\rangle=(0,1,0,0,0,0,0,0,0,0,0,0,0,0,0,0)$. The computational basis for a single qubit (two dimensions) is $|0\rangle=(1,0)$ and $|1\rangle=(0,1)$.

An alternative common basis consists of the eigenvectors of the Pauli-x operator: $\left.|+\rangle=\frac{1}{\sqrt{2}}(1,1)\right)$ and $\left.|-\rangle=\frac{1}{\sqrt{2}}(1,-1)\right)$.

So, for a classical state of $n$ bits, a $2^{n}$-dimensional probability distribution requires an exponential number of real numbers. We can think of the system as being one of the $n$-bit strings, but we do not know which one.

However, in quantum computing all $2^{n}$ complex coefficients need to be kept track of to see how the quantum computer calculates. For example, a 32 -qubit quantum computer has a state described by $2^{32}$ complex numbers.

## 4 Chromosome representation

The chromosome can be represented by the chromosome matrix, as follows [14]:

$$
Q=\left[\begin{array}{ccccc}
\alpha_{1} & \ldots & \alpha_{m} & \ldots & \alpha_{M}  \tag{3}\\
\beta_{1} & \ldots & \beta_{m} & \ldots & \beta_{M}
\end{array}\right]
$$

Moreover, the procedure of random selection of decision values is involved with a chromosome matrix. If the decision variable $x_{m}$ is characterized by ( $\alpha_{m}, \beta_{m}$ ), then is equal to 0 with the probability $\left|\alpha_{m}\right|^{2}$ and it is equal to 1 with $\left|\beta_{m}\right|^{2}$ [18].

AQMEA is working on a digital computer and collapsing into a single state does not occur in AQMEA. So, we simulate the process of observation.

A standard four-bit state and a quantum four-qubit state are sixteen-dimensional elements that are processed differently for standard or quantum computation.

In randomized computation, the application of stochastic matrices preserves that the sum of probabilities has to be equal to one. On the other hand, in quantum computation, allowed operations are unitary matrices. Those matrices are effectively rotations and preserve that the sum of the squares is equal to one [18].

A unitary matrix is an $n$ by $n$ complex matrix $Q$ satisfying the condition [18]:

$$
\begin{equation*}
Q^{*} Q=Q Q^{*}=I \tag{4}
\end{equation*}
$$

where
$Q^{*}$ - the conjugate transpose (the Hermitian adjoint) of $U$,
$I$ - the identity matrix in $n$ dimensions.
What sort of unitaries can be applied depends on the quantum devices.

Quantum computations are reversible because they are probabilistic combinations of unitaries. So, quantum computation generalize classical computation.

Upon termination of the algorithm, the result needs to be read off. In the case of a classical computer, we sample from the probability distribution on the four-bit register to obtain one definite four-bit string, e.g. 0100.

In quantum computation, we measure the four-qubit state, which is equivalent to collapsing the quantum state down to a classical distribution with the coefficients being the squared magnitudes of the coefficients for the quantum state. It is followed by sampling from that distribution. This destroys the original quantum state.

Many algorithms only give the correct answer with a certain probability; however by repeatedly initializing, running and measuring the quantum computer, the probability of getting the correct answer can be increased.

The binary chromosome $x=\left(x_{1}, \ldots, x_{m}, \ldots, x_{M}\right)$ can be measured with the probability $p(x)$ calculated, as follows:

$$
\begin{equation*}
p(x)=\prod_{m=1}^{M} p\left(x_{m}\right) \tag{5}
\end{equation*}
$$

where

$$
p\left(x_{m}\right)= \begin{cases}\alpha_{m}{ }^{2} & \text { for } x_{m}=0 \\ \beta_{m}{ }^{2} & \text { for } x_{m}=1\end{cases}
$$

Let us consider the chromosome, as below:

$$
Q=\left[\begin{array}{ccccc}
\frac{1}{\sqrt{3}} & \frac{1}{\sqrt{5}} & \frac{2}{\sqrt{7}} & \frac{1}{\sqrt{2}} & \frac{1}{2} \\
\frac{-\sqrt{2}}{\sqrt{3}} & \frac{2}{\sqrt{5}} & \frac{\sqrt{3}}{\sqrt{7}} & \frac{1}{\sqrt{2}} & \frac{\sqrt{3}}{2}
\end{array}\right]
$$

It means that the binary chromosome $(0,0,0,0,0)$ can be observed with the probability $1 / 210$, and the chromosome ( $0,0,0,0,1$ ) - $1 / 70$, and so on

Evolutionary computing with $Q$-bit representation has a better characteristic of population diversity than other representations, since it can represent linear superposition of states probabilistically. What is more, the search space The search space $\Omega$ can be generated by one Q -bit chromosome (however $\Omega$ consists of $2^{M}$ elements).

There is, as below:

$$
\begin{equation*}
\sum_{x \in \Omega} p(x)=1 . \tag{6}
\end{equation*}
$$

AQMEA is a probabilistic algorithm similar to other evolutionary algorithms. However, it maintains a population of $Q$-chromosomes. Although, the entire population is represented by one $Q$-chromosome, we consider the population of $\zeta Q$-chromosomes because of destroying the measured values. In fact, only one value of $Q$-chromosome can be measured and the others are supposed to be missed. We model this situation by the $\zeta$ $Q$-chromosomes based on complex numbers $\alpha_{m}$ and $\beta_{m}$, and formulas (2), (3).

An initial population can be generated by random numbers $\alpha_{m} \in[-1 ; 1]$ and calculation $\beta_{m}$ from (2). After
formulation the first $Q$-chromosome, the next one is formed until the whole population is created.

Then, we observe states of $Q$-chromosomes. A values of gene in the generated $x$-chromosome are selected either 0 or 1 for each bit using the probabilities from the related $Q$-chromosome. In a quantum computer, during the observing a quantum state, it collapses to a single state. However, collapsing into a single state does not occur in QEA, since it is working on a classical computer, not a quantum computer.

We observe the $Q$-chromosome from the population $\Theta(t)$, where $t$ is the number of population ( $t \leq T_{\max }$ ). We can observe the same chromosome $\lambda$ times. The parameter of sampling should be significantly smaller than size of the search space $2^{M}$, of course. In results, $\lambda x$-chromosomes are produced from one $Q$-chromosome. If the number of $Q$-individuals is equal to $\zeta$, the size of a binary population $L=\lambda \zeta$.

Afterwards, we evaluate the fitness for each binary chromosome by using the ranking procedure [2]. We determine non-dominated solutions from the current population and copy them to the archive.

Quantum approach for implementation of genetic algorithms has several advantages that are related to the previous applications. Integer factorization is computationally non-admissible by a classical computer for large integers that are the product of only a few prime numbers. On the other hand, a quantum computer could efficiently solve this problem using Shor's algorithm to find its factors. This ability would allow to "crack" many of the cryptographic systems, because there are a polynomial time algorithm for solving the problem. In particular, most of the public key ciphers are based on the difficulty of factoring integers or the related discrete logarithm problem, e.g. RSA. These are used to protect secure Web pages or encrypted email. It is a major result for electronic privacy and security. The approach to increase the security of an algorithm like RSA rely on increasing the key size to such amount that an enemy does not have a technology to build and use a powerful enough q-computer. However, it is possible to obtain the powerful quantum technology in the next decade.

More advanced approach is base on implementation quantum cryptography. There are some digital signature schemes that may be secured against quantum computers, e.g. Lamport signatures [18].

Quantum algorithms give not only polynomial speedup over the classical procedures, including the simulation of quantum physical processes from chemistry and solid state physics, the approximation of Jones polynomials, and solving Pell's equation [17]. For some questions, quantum algorithms give a polynomial speedup. An example is quantum database search that can be solved by Grover's algorithm using quadratically
fewer queries to the database than by standard computers. The plus is provable. Several other examples of provable quantum speedups for query problems are the finding collisions in two-to-one functions and evaluating NAND trees [26].

## 5 Genetic operations

In Adaptive Quantum-based Multi-objective Algorithm AQMEA, there are two populations. The first one is a $Q$-population $\Theta$ that consists on $\zeta Q$-individuals, and the second one is a $x$-population $P$ that consists on $L$ individuals. Moreover, each $Q$-chromosome generates $\lambda x$-chromosomes, randomly. It means that the quantum chromosome can be characterized by the selection probability $p_{s}\left(Q_{i}\right)$, as follows:

$$
\begin{equation*}
p_{s}\left(Q^{i}\right)=\frac{\sum_{j=1}^{\lambda} f\left(x^{i j}\right)}{\sum_{x^{i j} \in P(t)} f\left(x^{i j}\right)}, \quad i=\overline{1, \zeta} \tag{7}
\end{equation*}
$$

where
$Q^{i}$ - the $i$ th quantum chromosome from the current $Q$ population $\Theta(t)$,
$x^{i j}$ - the $j$ th binary chromosome determined randomly from the $i$ th quantum chromosome; $x^{i j}$ belongs to the current $x$-population $P(t)$.

We can prove, as follows:

$$
\begin{equation*}
\sum_{i=1}^{\zeta} p_{s}\left(Q^{i}\right)=1 \tag{8}
\end{equation*}
$$

A quantum crossover is based on the cross-overing of two matrixes given by formula (3). Each of matrixes is supposed to represent the $Q$-individual selected from the current quantum population with the probability $p_{s}$.

A short overview of classical evolutionary algorithms for multi-objective optimization problems is submitted in $[3,4]$. The name "adaptive evolutionary algorithm" for evolutionary algorithms is related to the changing of some parameters as a crossover probability, a mutation rate, a population size, and the others during the searching [5].

For considered algorithm, the crossover probability is decreased due to the number of new generations, as follows:

$$
\begin{equation*}
p_{c}=e^{-t / T_{\max }} \tag{9}
\end{equation*}
$$

where $e$ denotes the Euler constant.
Figure 1 shows a scheme of the adaptive quantumbased multi-criterion evolutionary algorithm AQMEA. We randomly generate $\zeta Q$-chromosomes for the quantum population $\Theta(0)$, where $\zeta \leq 10$. Because the
quantum algorithm is simulated on the classical von Neumann's computer, the structure of Q-chromosome is determined by (3) subject to (2). An initial population $\Theta(0)$ can be generated by random numbers $\alpha_{m} \in[-1 ; 1]$ and calculation $\beta_{m}$ from (2).

Then, we produce an initial binary population $P(0)$ that consists od $\lambda x$-chromosomes generated from each quantum-chromosome. If the gene $x_{m}$ is characterized by $\left(\alpha_{m}, \beta_{m}\right)$, then it is equal to 0 with the probability $\left|\alpha_{m}\right|^{2}$ and it is equal to 1 with $\left|\beta_{m}\right|^{2}$.

## 1. BEGIN

2. $t:=0, t$ - the number of population
3. set $\zeta$ the size of $Q$-population $\Theta, L$ size of binary population $\boldsymbol{P}, L=\lambda \zeta$, for the given sampling parameter $\lambda$
4. $p_{m}:=1 /(M \zeta), M-$ the length of $x$
5. generate an initial population $\Theta(0)$ and $\boldsymbol{P}(0)$,
6. calculate non-dominated ranks $r(x)$ and fitness
$f(x), x \in \boldsymbol{P}(t)$
7. finish:=FALSE
8. WHILE NOT finish DO
9. BEGIN $/ *$ new populations $\Theta$ and $\boldsymbol{P} * /$
10. $t:=t+1, \boldsymbol{P}(t):=\varnothing, \quad \Theta(t):=\varnothing$
11. calculate the selection probabilities $p_{s}(x), \boldsymbol{x} \in \boldsymbol{P}(t-1)$ by (6)
12. FOR $\zeta / 2 \mathrm{DO}$
13. BEGIN /* reproduction cycle */
14. 2 WT -selection of a potential parent pair $\{\boldsymbol{a}, \boldsymbol{b}\}$ from $\Theta(t-1)$
15. $Q$-crossover of a parent pair $\{\boldsymbol{a}, \boldsymbol{b}\}$ with the adaptive crossover rate $p_{c}, p_{c}:=e^{-t / T_{\max }}-0.1$
16. $Q$-mutation of an offspring pair $\left\{\boldsymbol{a}^{\prime}, \boldsymbol{b}^{\prime}\right\}$ with the adaptive mutation rate, $p_{m}:=\frac{t}{\zeta M T_{\text {max }}}$
17. $\Theta(t):=\Theta(t) \cup\left\{a^{\prime}, b^{\prime}\right\}$
18. END
19. generate $\boldsymbol{P}(t)$ by observing $\boldsymbol{\Theta}(t) \lambda$ times
20. calculate ranks $r(x)$ and fitness $f(x), x \in \boldsymbol{P}(t)$
21. IF $\left(\boldsymbol{P}(t)\right.$ converges OR $\left.t \geq T_{\max }\right)$ THEN
finish:=TRUE
22. END
23. END

Fig. 1. An adaptive quantum-based multi-criteria evolutionary algorithm AQMEA

The binary search space consists on $2^{M}$ elements that are represented by $x$-chromosomes. If $x$ is admissible, then the fitness function value is estimated, as below:

$$
\begin{equation*}
f(x)=r_{\max }-r(x)+P_{\max }+1 \tag{10}
\end{equation*}
$$

where
$r(x)$ - the rank of an admissible solution, $1 \leq r(x) \leq r_{\text {max }}$,
$P_{\text {max }}$ - the maximal value of the penalty function defined for non-admissible solutions.

In the quantum two-weight tournament selection (Fig. 1, line 14), the roulette rule is carried out twice due to $p_{s}\left(Q^{i}\right)$ for each $Q^{i}$ from the current quantum population. If two potential $Q$-parents $(a, b)$ are selected, we observe a $Q$-parent $\lambda$ times and produce $\lambda$ binary chromosomes. Then, we determine $N(a)$ - the set of non-dominated solutions from binary chromosomes generated for the quantum individual $a$. Similarly, we obtain $N(b)$. There is possible that one solution is included to the $N(a)$ or $N(b)$. In fact, it could be for the superior solution that dominates the others from the temporary set.

Afterwards, individuals from $N(a)$ and $N(b)$ are compared due to the Pareto relationship of domination. The dominated individuals are eliminated and $N(a, b)$ - the new non-dominated Pareto solution set is created. If at least one solution from $N(a)$ belongs to $N(a, b)$, then the quantum chromosome $a$ is accepted. The same rule is applied for the selection for cross-overing the $Q$ chromosome $b$.

If potential quantum parents $a, b$ generate binary solutions that are non-admissible, then the alternative with the corresponding smaller penalty is selected. Then, the random selection is repeated.

The fitness sharing technique can be substituted by the adaptive changing of main parameters and the generation of binary solution by observing quantum individuals. The quality of attained solutions increases in optimization problems with one criterion, if the crossover probability and the mutation rate are changed in an adaptive way.

The crossover point is randomly chosen for the quantum chromosome in the $Q$-crossover operator (Fig. 1 , line 15). The crossover point is selected between two columns of the matrix (3) and separates it on two submatrixes. These sub-matrixes are exchanged with corresponded sub-matrixes from the other quantum chromosome. It is worth to notice that crossover is carried out on quantum chromosomes instead of binary chromosomes as at classical genetic algorithm.

The crossover probability is equal to 0.9 at the initial population and almost each pair of potential parents is obligatory taken for the crossover procedure. A crossover operation supports the finding of a highquality solution area in the search space. It is important in the early search stage, especially. If the number of generation increases, the crossover probability decreases. Some search areas with the high quality solutions are identified after several crossover operations. That is why, value $p_{c}$ decreases to 0.5065 , if $t=100$ for
maximum number of population $T_{\max }=200$. The final smallest value $p_{c}$ is 0.2679 .

In $Q$-mutation (Fig. 1, line 16), the random swap of the column from matrix (3) by another one is applied. If the gene $\left[\begin{array}{l}\alpha_{m} \\ \beta_{m}\end{array}\right]$ is randomly taken for mutation, the new value $\alpha_{m}$ is randomly taken from the period $[-1 ; 1]$. Then, the complementary value $\beta_{m}$ is calculated from (2). A mutation rate increases according to the progress of the generation number.

To improve the quality of solution, we propose the development of the negative selection algorithm (NSA) from an immune systems. The immune system can be seen as a distributed adaptive system. The negative selection algorithm is based on the discrimination principle that is used to know what is a part of the immune system is.
An antigen is a molecule that stimulates a response against trespassers. The term originated from the notion that they can stimulate antibody generation. Moreover, the immune system consists of some viruses as well as bacteria. An antibody (an immunoglobulin) is a large $Y$-shaped protein used to identify and neutralize foreign objects like bacteria and viruses. The antibody recognizes a specific target. The negative selection can be used to manage constraints in an evolutionary algorithm by isolating the contemporary population in two groups. Feasible solutions called "antigens" create the first cluster, and the second cluster of individuals consists of "antibodies" - infeasible solutions.
We assume the initial fitness for antibodies is equal to zero. Then, a randomly chosen antigen $G^{-}$is compared to the selected antibodies. After that, the distance $S$ between $G^{-}$and the antibody $B^{-}$is calculated due to the amount of similarity at the genotype level. The measure of genotype similarity between the antigen and the antibody depends on their representation. This assessment of similarity for the integer version is, as follows [3]:

$$
\begin{equation*}
S\left(G^{-}, B^{-}\right)=\sum_{m=1}^{M}\left|G_{m}^{-}-B_{m}^{-}\right|, \tag{11}
\end{equation*}
$$

where
$M$ - the length of the solution,
$G_{m}^{-}$- value of the antigen at position $m, m=\overline{1, M}$,
$B_{m}^{-}-$value of the antibody at position $m, m=\overline{1, M}$;
The negative selection can be implemented by an external genetic algorithm to the AQMEA. In that approach, infeasible solutions that are similar to feasible ones are preferred in the current population. Although, almost all the random choices are based on the uniform distribution, the pressure is directed to improve the fitness of appropriate infeasible solutions.

## 6 Convergence of AQMEA

Let the Pareto points $\left\{P_{1}, P_{2}, \ldots, P_{U}\right\}$ be given for the considered instance of the optimization problem with $N$ criteria, and let points $\left\{A_{1}, A_{2}, \ldots, A_{W}\right\}$ be produced by an algorithm. The level of convergence to the Pareto front is calculated due to the Euclid distance, as follows:

$$
\begin{equation*}
S=\sum_{w=1}^{W} \sqrt{\sum_{n=1}^{N} \min _{u=1, U}\left(P_{u n}-A_{w n}\right)^{2}} . \tag{12}
\end{equation*}
$$

An average level $\bar{S}$ is calculated for several runs of the evolutionary algorithm.

As reported in [3], the best outcomes were obtained for some task assignment problems by the AMEA+. This algorithm gives better results than the previous AMEA. After 200 generations, an average level of Pareto set obtaining is $1.8 \%$ for the AMEA $+3.4 \%$ for the AMEA. 30 test preliminary populations were prepared, and each algorithm starts 30 times from these populations. For integer constrained coding of chromosomes, there are 12 decision variables and the search space consists of 25 600 solutions.

For the other instance with 15 tasks, 4 nodes, and 5 computer sorts there are 80 binary decision variables. An average level of convergence to the Pareto set is $16.7 \%$ for the AMEA+ and $18.4 \%$ for the AMEA. A maximal level is $28.5 \%$ for the AMEA+ and $29.6 \%$ for the AMEA. For this instance the average number of optimal solutions is $19.5 \%$ for AMEA+ and $21.1 \%$ for AMEA.

An average level of convergence to the Pareto set, an maximal level, and the average number of optimal solutions become worse, when the number of task, number of nodes, and number of computer types increase. An average level is $34.6 \%$ for the AMEA+ versus $35,7 \%$ for the AMEA, if the instance includes 50 tasks, 4 nodes, 5 computer types and also 220 binary decision variables.

It is worth to mention that those calculations have been carried out by a classic computer, but the expected results can be obtained for quantum computers. Consider a problem of a password cracker. In this question there is the only way to solve it by guessing answers repeatedly and check them. Moreover, there are some possible answers to check, and also every possible answer takes the same amount of time to check. Finally, there are no clues about which answers might be better: generating possibilities randomly is just as good as checking them in some special order [10]. In a password cracker problem, there are attempts to guess the password for an encrypted file. We assume that the password has a maximum possible length.

For such problems, the time for a quantum computer to solve this is proportional to the square root of $n$ (number of possible answers about password). That can be a very large speedup, reducing a calculation time
from months to milliseconds. It can be used to attack symmetric ciphers such as Triple DES and AES by attempting to guess the secret key [11].

Grover's algorithm can also be used to obtain a quadratic speed-up over a brute-force search for a class of problems known as NP-complete [10]. Since chemistry and nanotechnology rely on understanding quantum systems, and such systems are impossible to simulate in an efficient manner classically, many believe quantum simulation will be one of the most important applications of quantum computing [11].

## 7. Results for benchmark problem

To test the ability of the AQMEA, we consider a multicriterion optimization problem for task assignment in a distributed computer system, where three criteria are optimized. In the formulated task assignment problem as a multi-criterion question, both $Z_{\max }-$ a workload of a bottleneck computer and $C$ - the cost of system are minimized; in contrast, $R$ - a reliability of the distributed system is maximized. Moreover, there are constraints for the performance of the distributed systems and the probability that all tasks meet their deadlines. In addition, constraints related to memory limits and computer locations are imposed on the feasible task assignment.

The first criterion is the workload of the bottleneck computer for the allocation $x$, and its values are provided by the subsequent formula [5]:

$$
\begin{equation*}
Z_{\max }(x)=\max _{i \in \overline{1}, I}\left\{\sum_{j=1}^{J} \sum_{v=1}^{V} t_{v j} x_{v i}^{m} x_{i j}^{\pi}+\sum_{\left.\substack{v=1 \\ V} \sum_{\substack{u=1 \\ u \neq v}}^{V} \sum_{\substack{i=1 \\ k \neq i}}^{I} \sum_{k=1}^{I} \tau_{v u i k} x_{v i}^{m} x_{u k}^{m}\right\}}^{x}\right. \tag{13}
\end{equation*}
$$

where
$x=\left(x_{11}^{m}, \ldots, x_{1 I}^{m}, \ldots, x_{v i}^{m}, \ldots, x_{V I}^{m}, x_{11}^{\pi}, \ldots, x_{1 J}^{\pi}, \ldots, x_{i j}^{\pi}, \ldots, x_{I J}^{\pi}, N_{1}, \ldots, N_{v}, \ldots, N_{V}\right)$,
$\tau_{\text {vuik }}$ - the total communication time between the task $T_{v}$ assigned to the $i$ th node and the $T_{u}$ assigned to the $k$ th node.

Figure 2 shows three cuts in task assignment graph. We can balance workload among several processors by finding an optimal value of the bottleneck computer.

Let $\pi_{j}$ be failed independently due to an exponential distribution with rate $\tilde{\lambda}_{j}$. We do not take into account of repair and recovery times for failed computer in assessing the logical correctness of an allocation. Instead, we are supposed to allocate tasks to computers on which failures are least likely to occur during the execution of tasks. Computers and tasks can be assigned to nodes in purpose to maximize the third criterion - the reliability function $R$ defined, as below [4]:

$$
\begin{equation*}
R(x)=\prod_{v=1}^{V} \prod_{i=1}^{I} \prod_{j=1}^{J} \exp \left(-\tilde{\lambda}_{j} t_{v j} x_{v i}^{m} x_{i j}^{\pi}\right) \tag{14}
\end{equation*}
$$



Fig. 2. Load balancing by finding an optimal task assignment
The second measure of the task assignment is a cost of computers that is calculated, as below:

$$
\begin{equation*}
C(x)=\sum_{i=1}^{I} \sum_{j=1}^{J} \kappa_{j} x_{i j}^{\pi} \tag{15}
\end{equation*}
$$

where $\kappa_{j}$ corresponds to the cost of the computer $\pi_{j}$.
The minimal performance of the distributed systems $\Xi_{\min }$ is supposed to be smaller than the performance of the entire system that can be estimated according to the following formula:

$$
\begin{equation*}
\Xi(x)=\sum_{i=1}^{I} \sum_{j=1}^{J} \delta_{j} x_{i j}^{\pi} \tag{16}
\end{equation*}
$$

where $\delta_{j}$ is the numerical performance of the computer $\pi_{j}$ for the task benchmark, for instance [MFlops].

The probability that all tasks meet their deadlines is supposed to be greater than the minimal probability $P_{\min }$. This parameter is usually set to be greater than 0.9.

$$
\begin{equation*}
P_{D}(x)=\sum_{l=1}^{K} p_{i} \prod_{m_{v} \in M_{l}} \xi\left(d_{v}-C_{v}(x)\right) \tag{17}
\end{equation*}
$$

Two main constraint types: the benchmark performance limit and also probability that all tasks meet their deadlines are supposed to be complement with some resource constraint.

Figure 3 shows the cut of the evaluation space that is explored by the most effective meta-heuristic AMEA* [3]. Evolutionary algorithm AMEA* [3], the ant
algorithm [12] and genetic programming MGP [4] have been applied for solving some versions of multi-criterion task assignment. We can compare quality of obtained solutions by AQMEA to qualities produced by the other multi-criterion meta-heuristics.

The binary search space consisted of $1.0737 \times 10^{9}$ elements and included 25600 admissible solutions. An average level $\bar{S}$ was calculated for fifty runs of the algorithm. After 350 assessments of those functions, an average level of Pareto set obtaining is $1.5 \%$ for the AQMEA, $1.7 \%$ for the MOTB.


Fig. 3. Pareto front and results of AMEA*
Quantum search provides a promising alternative for the other problems like the face verification problem [15], disk allocation [16] or the Travelling Salesman Problem [21].

There are a number of practical difficulties in building a quantum computer, and quantum computers have only solved trivial problems. David DiVincenzo listed some requirements for a practical quantum computer [11]. Quantum computers are supposed to be scalable physically to increase the number of qubits. Qubits should be initialized to arbitrary values and quantum gates are expected to perform faster than decoherence time. What is more, a universal gate should be set, and qubits are supposed to be read easily.

A crucial question is controlling or removing decoherence what is related to isolating the system from its environment as the smallest interaction with the external systems would cause the inner system to de-cohere. This result is irreversible, as it is non-unitary, and is generally something that is supposed to be avoided, if not highly
controlled. The transverse relaxation (de-phasing) time for some technologies, typically range between nanoseconds and seconds at low temperature.

These implementations are more complicated for optical approaches when the timescales are orders of magnitude lower and an optical pulse shaping is applied. Error rates are usually proportional to the proportion of operating time to de-coherence time. So, an operation is required to be completed much more quickly than the de-coherence time [18].

If the error tempo is small enough, it is possible to apply quantum error correction, which corrects errors due to de-coherence. This is the constraint that the total calculation time should be longer than de-coherence period. This implies that a gate is supposed be able to execute its job $10^{4}$ times faster than the de-coherence time of the computer [10].

The development of error correction is related to the increased number of required qubits. The number required to factor integers using Shor's algorithm is between $n$ and $n^{2}$, where $n$ is the number of bits in the number to be factored. Error correction procedure inflates this result by an additional factor of $n$. For $n=100$, it implies necessitate for about $10^{3}$ qubits without error correction. With error correction, the figure would rise to about $10^{5}$ qubits [11].

Another approach to the stability-decoherence problem is to create a topological quantum computer with anyons, quasi-particles used as threads and relying on braid theory to form stable logic gates [18].

In 2009, researchers at Yale University created the first rudimentary solid-state quantum processor. The two-qubit superconducting chip is able to run elementary algorithms. Each of the two artificial qubits are made up of a billion aluminum atoms but they acted like a single one that could occupy two different energy states [10].

## 8. Concluding remarks

To find optimal solutions, the quantum-based adaptive evolutionary algorithm AQMEA is proposed. It is an advanced technique for finding Pareto-optimal task allocation problem with the maximization of the system reliability and distributed system performance. Moreover, the workload of the bottleneck computer and the cost of computers are minimized.

Although, there are several technological problems in building a quantum computer, quantum search provides a promising alternative for development some genetic algorithms and others multi-objective optimization techniques.

Our future works will concern on a development the combination between quantum computing and evolutionary algorithms for finding Pareto-optimal solutions.

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