The quantum language of the microRNA gene and anticancer: With a dynamic computer simulation of human breast cancer drug resistance

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Abstract

Precision medicine requires for treatments according to the personalized genetic and environmental diversities. In insights of the diversity, individual variation of drug sensitivity would be implied in many factors, one of them would be the microRNA (miRNA) gene. The drug resistance would also be related with alteration of miRNA genotype in the cancer cell via its administered anticancer drug. Further, natural products as the nutraceuticals could also modulate miRNA expression and promote anticancer effects. In the case of foods, not only foods' miRNAs but also nutrients and natural products could render changing the phenotype of cells by alteration of miRNA expression. The facts show that environmental quantum energy of therapeutic drugs and foods may have an important role for miRNA gene expression upon the cells to personalize biological reaction under pharmacokinetics, and the new pharmacokinetics may be required through miRNA response. In turn, drugs among individuals treated would be also affected by personal state of quantum energy in the miRNA genes, and the quantum language of miRNA would dynamically adjust re-position of the personalized health. On the contrary, miRNA could control protein expression or epigenetic traits both negatively and positively, or directly and indirectly in the transcription and translation processes. Dysregulation of miRNA induces human cancer. Further, long noncoding RNA (lncRNA) and circular RNA (circRNA) could also act together with miRNAs and would control protein gene expression with miRNA. Participating in cancer development and progression, lncRNAs and circRNAs would directly or indirectly regulate signalling pathways and proliferation processes via miRNA sponging, and function as miRNA reservoir or as binding protein scavengers. The pharmaceutical agents would affect these RNA gene regulators at first in a cell and in vivo. To elucidate new pharmacokinetics of agents, the miRNA-miRNA-lncRNA-circRNA network architecture should be investigated. Thus, we review lncRNA and circRNA functions in cancers. Then, quantum energy relation among miRNA-miRNA-lncRNA-circRNA is discussed. Further, as a sample of drug treatment-related with quantum language of miRNA, drug resistance of human breast cancer was dynamically simulated using the miRNA entangling target sorter (METS). Since we have proved that quantum characters among miRNAs is implicated in oncogenesis, experimental evidences reviewed and dynamic in silico simulation with METS suggested that quantum language of miRNAs may be a common factor through tumorigenesis, anti-cancer and drug resistance.

Overview

While >90% of the human genome is transcribed, the dominant transcripts are noncoding RNA [1]. ncRNAs are classified as housekeeper and regulator, however, recently housekeepers of ribosome RNA (rRNA) and transfer RNA (tRNA) have been shown that they would produce miRNAs and transfer RNA fragments as the regulator [2,3]. Therefore, including long noncoding RNA (lncRNA) and circular RNA (circRNA), ncRNAs would be the regulators of protein expression [1,4-6]. miRNA is the noncoding gene while lncRNA is epigenetic regulator, but lncRNA also sometimes contains coding region or antisense of protein gene, further it remains forever so that the lncRNA is a source of the resident miRNA genes [7,8]. As described below, it seems to be doubtful what it is epi-gene because protein-coding DNA is absolutely transcribed to mRNAs and lncRNAs, which are also transcribed from protein-noncoding DNA, so it contains the protein gene. Thus, here, we define lncRNA and circRNA as two of the noncoding genes. These noncoding RNA (ncRNA) genes are implicated in treatment of chemotherapeutic agents, in diet as well as in immunotherapeutic agents. The new pharmacokinetic era has recently been revealed by miRNA-lncRNA-circRNA network (Figure 1). In addition, while miRNA dysregulation in drug resistance of human breast cancer has been reviewed from several independent studies, using the miRNA entangling target sorter (METS), breast cancer drug resistance was, in silico, simulated for preparation of the personalized precision medicine [9].

Long noncoding RNA

LncRNA is also categorized as the regulator according to >200 nts long [8]. LncRNA class contains four types of the genomic organization (Figure 2).

Intergenic lncRNAs are derived from transcripts among protein genes. Intronic lncRNA is from introns. lncRNA overlapped exon and as antisense are the third and fourth classes. LncRNA acts as a transcriptional signal on one to one of Pol II, molecular decays, guide of protein gene expression in cis and trans, and the scaffold molecule [4]. However, the four archetypes did not contain the relation to miRNAs, therefore, it has been explained that lncRNA, such as Air mapped in imprinting gene cluster Ig2r and would repress histone modification in allele-specific manner but in the imprinting site, methyltransferase
Factors of forehead box C1 (FOXC1) and zonula occludens 2 (ZO-2) by increasing expression of miR-137 [18]. PVT1 overexpression in glioma vascular endothelial cells induced protective autophagy by upregulation of Atg7 and Beclin 1, which would be targeted by miR-186 [19]. PVT1 could be a sponge of miR-186. Further, HOXAs2 was increased in glioma cells [20]. HOXAs2 bound to miR-373 and miR-373 targets to epidermal growth factor receptor (EGFR). Since glioma malignancy is deeply implicated in vascularization process via EGFR signalling pathway, such as vascular endothelial-cadherin (V-cadherin), matrix metalloproteinase-2 (MMP-2) and -9 (MMP-9) through phosphatidylinositol 3 (PI3)- kinase pathway, HOXAs2/miR-373 interaction showed the relation of a molecular sponge. HOX transcript antisense RNA (HOTAIR) lncRNA expression is inversely related with miR-148b-3p in glioma cells, miR-141 as well [21,22].

miR-34a/b/c has deeply been implicated in anti-tumour ability of p53 against wide spectrum of cancers [7]. LncRNA, LINC00473 was targeted by miR-34a in cervical cancer, Lnc4-c blocked miR-34a to bind miR-34a encoded gene in colon cancer stem cells [23,24]. About drug resistance of cancer cells, KO of LncRNA HOTAIR suppressed cisplatin resistance of gastric cancer by upregulation of miR-34a expression [25]. HOTAIR is a sponge of miR-34a. A soy isoflavone, genistein inhibited prostate cancer cells via reducing HOTAIR, resulting up-regulation of miR-34a [26]. It is suggested that natural product would be targeting to LncRNA and miRNA expression, probably through relation of quantum energy among chemical substances. In etiology of cancers, LncRNAs are implicated in cancers, such as breast cancer, however, the relation among miRNAs and LncRNAs has not been enough shown to simulate the network of miRNA/LncRNA/mRNA [27].

LncRNA would be indirectly regulated by protein expression through miRNAs. These data suggested that a lncRNA could be targeted by multiple miRNAs and a miRNA would be sponged by multiple lncRNAs. Those relations would be regulated through the similar rule to the implication between miRNAs and mRNAs. It means that mRNA as well as IncRNA would usually not be cleaved by the targeting of miRNAs and would functionally suppress to mRNA vice versa.

Circular RNA (circRNA)

Circular RNA (circRNA) is belong to noncoding RNA (ncRNA) and is bio-generated from thousands of protein coding exon and intron (Figure 3).

The dominant circRNAs are cytoplasmic 3'→5'-linked ones consisting of exon (EcircRNA). The nuclear localized circRNAs are 2'→5'-linked ones consisting of intron (circRNA). Spliceosome-dependent mechanisms lead to formation of circRNAs. Conventional collinear splicing causes the excision of intron from a multi-exon of protein gene and ciRNA is produced. Another nuclear localized circRNAs are 3'→5'-linked ones consisting of both exon and intron (EciRNA). The 3'→5'-linked EciRNAs are formed by back splicing. The back splicing is required for location of the 3'-end of an exon in closely proximity to an upstream 5'-end of the same or another exon. The looping structure by back splicing would be facilitated by the flanking reverse complimentary Alu retrotransposons or dimerizing RNA binding proteins. Further, circRNA could be bio-generated by an exon containing lariat from an exon-skipping events. The ciRNAs and EciRNAs are secreted from cells and presented in blood, body fluids and tissues [28]. The function of circRNA against RNA gene information is implicated in the sponge effects to miRNAs and ciRNAs would be a source of intronic miRNAs as mitrons and simtrons [7]. Since LTR-retrotransposon, Alu and LINE have circular shape, and
they have pre-miRNAs and miRNA target sequences, circRNA as well as the retroelements would facilitate the RNA crosstalk with miRNAs in the same functions. Since circRNA is bio-generated by splicing and the splicing is controlled by miRNAs, circRNA would simply have been a target of miRNA [29]. Although circRNA has other functions, such as stimulation of the initiation and elongation of RNA polymerase II transcripts by ElcIRNA, RNA binding protein reservoir and translation of an exon including N (6)-methyl adenosine-driven translation of circRNA, these transcriptional and translational regulations would be restricted by canonical mRNA transcription and translation [28].

Therefore, circRNA and miRNA interaction would have more important role for protein gene expression to precisely control life events, such as cell proliferation and development than circRNA and protein direct interaction. In the competing endogenous RNA (ceRNA) theory, miRNAs could compete against target elements of mRNAs, long noncoding RNAs (lncRNAs) and circRNAs in the cytoplasm at the same time. If so, gain and loss effects of miRNA information could not be obtained from simple experiments on the bench. However, large amounts of research on the bench have showed that single miRNA gain and loss experiments affected the expression of target proteins. Thus, ceRNA is just the system but not language as information [30]. On the contrary, absolutely miRNA-miRNA quantum code would be information one [31].

**Network computing analysis**

To investigate the etiology of cancers from big data base using miRNA biomarkers, network analysis would be an important tool. The core networks and hubs of cancers have been reported by several research groups. Although the experimental data would have not been enough to complete molecular mechanisms of cancers, the network computing analysis revealed complex relation among miRNAs and target mRNAs [32]. About 34 altered miRNA expression was predicted in the glioma Cpg island methylator phenotype (CIMP) (miR-19a/b, -1, -124, -206, -130a/b, -301, -29a/b/c, -494, -133a/b, -34/a/b/c, -449, -15a/b, -16, -186, -195, -424, -497, -140, -9, -302b, -145, -519a/b/c, -24, -506, and -122a). Further, glioblastoma has been investigated in IncRNA-mediated network prediction [33]. Chiu et al. has established the platform for 'Cupid' for computer inferring network of miRNA-miRNA and IncRNA [34]. Cupid using machine learning prediction was trained on biochemical assay data, therefore, network analysis has been closed to be physiologically relevant at least. While retrotransposons are regulated by miRNA so that it is not sponge, expression of IncRNA is also controlled by miRNAs [35]. Meanwhile, miRNA target sites have also been proved themselves to locate in the protein coding RNA sequences, such as exon. Piwecka et al. have shown that knockout (KO) of Cdr1as/cirs-7 circRNA in mouse brain causes depression of miR-7 and dysfunctional synaptic transmission [36]. Since miR-7 targets more than 70 binding sites of Cdr1as circRNA, exon, even noncoding and antisense, is actual target sites and miRNA did not break out the target of a sponge. Cdr1as would be degraded by miR-671 in Argonaute 2 (Ago2) dependent manner; however, Cdr1as was not cleared by miR-7 [37]. KO of Cdr1as resulted decreasing of miR-7 but not increasing, whose results are clearly supported by mathematical model as following description. Further, knockdown of IncRNA ACT104 in osteosarcoma cells inhibited their proliferation, migration and invasion by upregulation of tumour suppressor miR-381 [38]. Thus, it is suggested that miRNA is still an axis of ncRNAs' function because most of all proteins are directly controlled by miRNAs, and miRNAs would be tuned by other ncRNAs. Although we have been shown that HIV-1 N367 miRNA has the target site in various retroviruses and coding region has the target sites for N367, and RNA-derived miRNAs have target sites in transposable element (TE) LINE, miRNAs would have physiochemically increased target sites according to ceRNA theory [2,30,39]. By simplified mathematical formula of quantum computing, it will be cleared. Tensor products of miRNA (X), mRNA (P), IncRNA (Q), circRNA (R) and TE (S) are shown in the qubits as previously described [31].

\[
|\Psi_{x} > \otimes |\Psi_{y} > = |\Psi_{xy} > \\
|\Psi_{x} > \otimes |\Psi_{q} > = |\Psi_{xq} > \\
|\Psi_{x} > \otimes |\Psi_{r} > = |\Psi_{xr} > \\
|\Psi_{x} > \otimes |\Psi_{s} > = |\Psi_{xs} >
\]

Tensor products of X, P, Q, R and S in a dimension are also presented as follows,

\[
|\Psi_{x} > \otimes |\Psi_{y} > \otimes |\Psi_{q} > \otimes |\Psi_{r} > \otimes |\Psi_{s} > = |\Psi >
\]

Therefore, when i is 1, --------, n,

\[
|f(x) > ,
\]

the total quantum qubit is linearly transformed with the vector spaces of Xi, Pi, Qi, Ri and Si.

\[
|f(x) > = a_{0} \sum_{i=1}^{n} Xi Pi Qi Ri Si
\]

a_{0} is amplified value.

Xi, Pi, Qi, Ri and Si are the same formula as targets or sponges of miRNAs, therefore,

\[
|f(x) > = a_{0} \sum_{i=1}^{n} Xi Ti
\]

This formula is the same as that of DNSs among miRNAs [40]. From this formula, miRNA quantum energy can be accumulated; and it is strongly supported that plant or milk miRNAs are accumulated in transmitted organs and plant miRNAs accumulate in human breast milk [41-44]. The cumulative effects have showed in quantum language simulation, such as miRNA entangling target sorting (METS) [45]. Since miRNA DNSs are correlated with CSCs, quantum energy between miRNA and mRNA depends upon DNS of miRNAs [46]. Although expression of the miRNA genes would genetically control upon the expression rates of miRNAs, IncRNAs, circRNAs and TE, miRNA
expression could be controlled by miRNA itself in cancer [47]. By using the Cancer Genome Atlas (TCGA) data sets, the lncRNA-miRNA-miRNA network analyses have showed significant axis correlated with prognostic survival rate in lung squamous cell carcinoma [48]. The axis of network was PLAU mRNA, miR-31-5p, miR-453-3p, and lncRNAs, FAM83A-AS1, MIR31HG, and MIR99AHG. MIR31HG and MIR99AHG contain miR-31 and miR-99a host genes, respectively. Namely, since lncRNA, circRNA and TE are a source of pre-miRNAs and mature miRNAs, the increase-decrease rate would be involved in approximate parallel to that of miRNAs. Analyses of network among IncRNA, miRNA and mRNA have been performed in several cancers, such as pancreatic cancer, oesophageal cancer, hepatocellular carcinoma, colon cancer, rectal adenocarcinoma, gastric cancer, and thyroid carcinoma [49-55]. These data suggest that multiple miRNAs are the key driver to control miRNAs for cancer prognosis in the network.

Altogether, the resident miRNAs‘ package would be stored with sponge ncRNAs and as reservoir ncRNA, and miRNA/miRNA interaction is more important for target selection than previous ceRNA [45]. Further, its sponge function shows the clear evidence that miRNAs are usually not degraded but the quantum energy of miRNAs are stored.  

**Pharmaceutical effects through ncRNAs**

Chemotherapeutic agents react target proteins in vitro and it has been believed that such agents could work the same pharmacological activity in vivo as in vitro (Figure 1). Since first impact of agent transferred through the plasma membrane would be a chance meeting of large number of ncRNAs in the cytosol, miRNAs and IncRNA altered expression and tuned cell functions according to the response of ncRNAs against agent. Further, enzymes related with drug-metabolizing and transport, such as CYP450s and ABC or SLC transporters and xenobiotic receptors, could be controlled by the miRNA genes: therefore, if there were SNPs in miRNAs or in miRNA target sites, these genetic mutations are associated with chemotherapy response and clinical output [56]. It is suggesting that another pharmacokinetic era would be newly opened with the miRNA genes as well as their targets of IncRNA (Figure 1).

**Natural products**

Bufalin is a cardiotonic steroid from the Chinese medicine ChanSu. Treatment of Bufalin increased miR-203 expression and inhibition the stem cell-like phenotyping and induced apoptosis in glioma cells [57]. Bufalin prevented osteosarcoma cells through miR-148a [58]. Antipryretic analgesics in popular medicine, aspirin was correlated with expression of miR-145 in vascular smooth muscle cells, and its effects of anti-proliferation and anti-inflammation by aspirin were implicated in upregulation of miR-145 [59]. The omega-3 polyunsaturated acid (n-3 PFA), docosahexaenoic acid (DHA) reduced miR-21 oncomir in breast cancer because tamoxifen is a prodrug, which is bioactivated to endoxifen by CYP2D6 and CYP3A4 (Figure 4). Anti-HER2 human monoclonal antibody, trastuzumab is used for HER2 positive breast cancer. Triple-negative breast cancer did not have any target for its therapy, but doxorubicin would charge topoisoerase II and acts as DNA intercalator to inhibit DNA polymerization in breast cancer cells.

It is well known that ERα-positive, HER2-positive and triple negative breast cancer cells alter miRNA expression profile by treatment resistances of tamoxifen, trastuzumab and doxorubicin, respectively [71-76]. About etiological mechanisms from protein biology, tamoxifen resistant is implicated in phosphorylation of the serine 305 residue on ERα by protein kinase A (PKA) and p21-activated kinase-1. The 305-phosphorylation of ERα changes tamoxifen activity from antagonist of ERα to agonist but this mechanism is not fully cleared because endoxifen (tamoxifen) itself has no change binding to ERα and without chemical modification in culture [77]. In the microRNA gene and anti-cancer: With a dynamic computer simulation of human breast cancer drug resistance

Figure 4. Human breast cancer subtype. Simple markers of human breast cancer are represented. Percentage of phenotypes are different between data of researchers, therefore, approximate percentages are shown. Tamoxifen, trastuzumab (Herceptin) and doxorubicin (Adriamycin) are administered to ESR-positive, HER2-positive and triple negative breast cancer patients, respectively.
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general, signal transduction pathways are positively related with gene expression but not negative regulation. Although AKT2-activated ERα (phosphorylation) contributes to tamoxifen resistance, PI3K binds together with ERα, and subsequently AKT2 and PI3K (PKC pathway far from PKA) activates ERα by a non-transcriptional and ligand-independent mechanism in vivo and in vitro [78]. But Sun et al. has not described about relation between PI3K/AKT2 and tamoxifen resistance of breast cancer [78]. ERα is also activated by growth factors, such as fibroblast growth factor 1 and 3 (FGFR 1 and 3) and insulin-like growth factor receptor 1 (IGF-1R). These growth factors are implicated in tamoxifen resistance via PI3K/AKT signalling [79,80]. The mechanisms of trastuzumab resistance are epitope asking, signalling alteration and immune response by monoclonal antibody Fcγ receptor polymorphism. However, mechanisms of tamoxifen, trastuzumab and doxorubicin resistance have not yet been understood at all. To further investigation of drug resistance, the role of ncRNAs have been reported in breast cancer. Trastuzumab-responsible miRNAs were observed in trastuzumab-resistant breast cancer patients (HER-2 positive breast cancer) [81]. miR-210 was upregulated in trastuzumab-resistant BT474 cells in vitro. On the other hand, trastuzumab treatment of BT474 decreased miR-194 expression [82]. In the culture of trastuzumab resistant breast cancer cell, SKBr-3, miR-200c was significantly reduced [83]. Further, miR-129-5p was downregulated in trastuzumab-resistant human breast cancer cells (HER-2 positive breast cancer), JIMT-1 in vitro and in sera of patients [84]. Overexpression of miR-129-5p increased sensitivity to trastuzumab in JIMT-1. In the case of HER-2 positive gastric cancer, upregulation of miR-125b was significantly associated with trastuzumab resistance of HER-2 positive gastric cancer and was implicated in malignant progression and poor prognosis [85]. These data suggest that protein agents could affect expression of miRNAs in treated cancer cells and antibody agent has new pharmacokinetic activity to modulate miRNA expression. LncRNA ATB was upregulated in trastuzumab resistant SKBr-3 cells, and lncRNA GASS was downregulated in the same cell line and tissues from trastuzumab-treated patients [86,87]. The former was the sponge of miR-200c and the latter was that of miR-21. Chemotherapeutic anti-EGFR, lapatinib upregulated GASS in trastuzumab resistant SKBr-3, suggesting that functionally different agents show pharmacologically distinct effects against lncRNAs [87]. On the contrary, miR-21 increasing was correlated with treatment of both trastuzumab and anthracycline- or taxane-based chemotherapy in HER-2 positive breast cancer [88]. Therefore, responses of lncRNA GASS and miR-21 were different against each treatment in breast cancer at all. Further, since miR-21 would work as a gene but lncRNA would be a target, in actual, two RNAs would have been given upon the distinct function because the proof of concept for the miRNA gene has recently been finally confirmed in conserved miRNA loci of the inbred mouse strains‘ genome [89]. Although as described above, miRNA-miRNA interaction is important, Cilek et al. have simulated in the network analysis that trastuzumab-responsible miRNA-miRNA network has strong responsible for drug response in SKBr3 and BT474 breast cancer cells [90]. They found that miR-3064-3p and miR-32-3p were deeply connected to miR-216b and these miRNAs target YWHAE, RPL37 and AK2, which are related with apoptosis, cell cycle and metabolic pathway. Although alteration of miRNAs has been reported in trastuzumab resistance, trastuzumab treatment alters miRNA expression in HER2-positive breast cancer and would result in the resistant state [91]. We selected miRNAs for the MMP of trastuzumab resistance as downregulation of miR-200c, miR-221 and miR-205-5p, and upregulation of miR-375, miR-7-5p, miR-542-3p, miR-21, miR-210 and miR-515 (Table 1). Finally, pathway of antibody-receptor was regulated by miRNAs, therefore, drug affects expression of miRNAs directly and indirectly that is far from in vitro pharmacokinetic data, and the activity of drug in vivo cannot be correctly measured in vitro. Targets of trastuzumab resistance are mainly EGFR or PIK3/AKT in miRNA researches (Table 1), which shows an agreement with previous description about protein biology.

Simulation models of tamoxifen, trastuzumab and doxorubicin resistances

To elucidate the pathophysiological interactions between miRNAs and human breast cancer for personalized medicine, etiological causes of drug resistance were investigated by dynamic computer simulation in human breast cancer and its resistance according to systemic treatment. Since profiles of miRNAs have been available for prediction of human diseases, the microRNA memory packages (MMPs) were applied for the prediction of difference in drug resistance [46]. About in vivo trastuzumab resistance in breast cancer, 7 miRNAs were used, and it was compared with doxorubicin and tamoxifen resistances (Figure 5), in another word, at first contents of drug resistance were simulated before the context.

Backbones of drug resistant MMPs are ERα positive, HER2 positive and triple negative breast cancer ones. All MMPs were showed as unique shapes, and 6 types (three inputs and three outputs) of MMPs were obtained (Figure 5). Although human breast cancer is a heterogeneous disease, it seems that drug resistance would also have a heterogeneous phenotype. However, it is possible to be diagnosed by miRNA comb about prediction of drug resistance of human breast cancer, and it is strongly supported by which each MMP is unique. As artificial intelligence (AI) is programmed by deep learning algorithm into the layer architecture (Figure S1), therefore, we firstly screened drug resistant-related miRNAs in both downregulation (Figure S2 blue characters) and upregulation (Figure S2 red characters) to pick up from the PubMed and Google Scholar databases.

From matrix of DNSs in SNSs of selected miRNAs, layer structures were constructed depending on the quantum energy levels (Figure S3)

Table 1. Trastuzumab resistant miRNAs in breast cancer

| Drug  | miRNA | Target    | Mechanism                | SNS  | Reference number |
|-------|-------|-----------|--------------------------|------|------------------|
| Trastuzumab | **miR-375** | IGFR, AKT   | Activate PI3K/Akt pathway | 7    | 104              |
|       | *miR-7-5p  | EGFR       | Enhance EGFR expression  | 7    | 105              |
|       | *miR-200c | ZNF17      | Promote TGF-beta signalling | 6    | 83               |
|       | **miR-542-3p** | AKT       | Activate PI3K/Akt pathway | 5    | 125              |
|       | **miR-21** | PTEN, PDCD4 | Activate PI3K/Akt pathway | 5    | 126              |
|       | *miR-221  | PTEN       | Activate PI3K/Akt pathway | 4    | 73               |
|       | **miR-210** | MET, IGFR1 | Tumour growth           | 4    | 81               |
|       | *miR-205-5p| P63        | Promote EGFR pathway     | 4    | 127              |
|       | **miR-515** | MARK4      | Suppression PIK3/MARK4   | 3    | 128              |

*: downregulation, **: upregulation

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and depicted as mountain views (Figure S4). Three-dimension (3D) map was represented according to 10 DNS contour lines about DNSs of trastuzumab, doxorubicin and tamoxifen resistant-related miRNAs.

Next, context of drug resistance was simulated to understand difference of MRP. Since the network related with cancer has been dynamically computed by microRNA entangling target sorting (METS) algorithm, the network was predicted by METS simulator about \textit{in vivo} ER\textalpha positve breast cancer and it was compared with the network of tamoxifen resistance [45]. DNSs of 9 miRNAs in the matrix of tamoxifen resistance were calculated and top 4 values of DNS (70-80 called as quantum code region, QCR) were selected as miR-375, miR-342-5p, miR-320a and miR-378a-3p. The multi-targets of four miRNAs were searched in miRTarBase V21 (miRTarbase.mbc.nctu.edu.tw/index.php) and TargetScan 7.2 (www.targetscan.org/vert_72/) in human, then, target genes and related miRNAs were simulated by METS. The multi-miRNAs in a protein mRNA target were also reversely selected according to the top 10 DNS. Top layer (QCR: 70-80) of tamoxifen-resistant-related miRNAs were compared with the same layer (QCR: 70-80) of ER\textalpha positive breast cancer in target protein-protein interaction and the cluster on String (https://string-db.org/cgi/input.pl) (Figure 6A). Finally, protein gene function was searched by GeneCards (http://www.geneCards.org/) and validation of statistical significance in the simulation, such as the area under the ROC curve (AUC) or the \chi^2-based Cochran’s Q test was computed by BellCurve for Excel (Social Survey Research Information Co. Ltd., Tokyo, Japan). About searching of lncRNA, Long Noncoding RNA Database was used (www.lncrnadb.org). In ER\textalpha positive breast cancer (Figure 6a), three core miRNAs, let-7a-5p, miR-34a-5p and miR-206 are downregulated. Under METS simulation (Figure 6b), ER\alpha would be upregulated by miR-206 and miR-18a-5p (miR-17-92 cluster) downregulation as tumour suppressors of breast cancer [92-94]. Downregulation of miR-34a-5p contributes suppression of TP53 in breast cancer [95,96]. Decreasing let-7a (let-7 family) and miR-34a is correlated with anti-apoptotic pathway [97]. After resistance of tamoxifen treatment, miR-375 downregulation would increase PI3K expression (Figure 6c), which has induced activation of ER\alpha together with AKT as described above. As shown in GO enrichment analysis (geneontology.org/page/go-enrichment-analysis) (Figure 6c), cytoplasmic region would be PI3K. On the contrary, targets of miR-206 and let-7 family were changed from ER\alpha, and apoptosis-related transcriptional regulator, high mobility group A1 (HMGA1) to cytoplasmic vesicle-related complex 2 (CPLX2) and chaperon protein, translocase of inner mitochondrial membrane (TIMM), respectively. These simulations strongly supported that alteration of ER\alpha phosphorylation from PKA to PKC pathways is dominantly implicated in tamoxifen resistance in human breast cancer.

In the case of HER2 positive breast cancer, the networks of QCR 90-100 and QCR 40-50 were investigated at first, and then those of HER2 positive and trastuzumab resistant resistance were compared at the same QCR-levelled layers (QCR 40-50) by METS (Figure 7a). As shown in Figure 7b, downregulation of miR-134-5p, miR-637 and miR-193a-5p would induce upregulation of STAT, nuclear factor I/C (NFIC) and ribosomal protein S2 (RPS2). STAT and NFIC are transcriptional activator via HER2 signalling and RPS2 is a component of the 40S subunit, respectively. RPS2 is associated with Diamond-Blackfan Anaemia as ribosomopathy and activates mRNA production via formation of the cap-binding complex and eIFs [2]. Further, these three miRNAs and their comb miRNAs would cooperate with regulation of mTOR, PI3K and KRAS. The HER is belong to the epidermal factor receptor (EGFR) tyrosine kinase receptor family and amplification of it stimulates MAPK, PI3K and mTOR pathways [98]. This evidence is equivalent to our prediction by METS.

In the layer of QCR 40-50 (Figure 7c), downregulation of miR-498, human specific miR-1273d and miR-1295b-3p are linked in RPL37A, which is a component of 60S subunit of ribosomal complex. Therefore, HER2 positive human breast cancer would be involved into a case of ribosomopathy and this results from simulation would be a new insight of etiology in HER2 positive human breast cancer. Human breast cancer has been implicated in ribosomopathy, however, etiological causes of dysregulation of ribosomal proteins have not yet been explained with their gene- or oncoprotein gene- and tumour suppressor protein gene amplification, deletion, mutation and SNPs in the data from TCGA [99]. Thus, it is suggested that dysregulation of miRNAs could induce ribosomopathy, and anti-ribosomopathy would target development of agents against human HER2-positive breast cancer. Furthermore, miR-498 and its comb miRNAs are implicated in tumour suppressor, BCC1A, and miR-542-5p would target EGFR. After treatment of trastuzumab, most of patients respond trastuzumab resistance within a year [100]. As shown in Figure 7d, miR-375 can target SP1 and miR-7-5p targets IGF1R. ABCC1, BCL2, and EGFR and SP1 are connected as cell membrane signalling and tumour proliferation. To validate the prediction by METS, dominant relations (red circles) were applied for GO cellular component analyses (Figure 7d). It is predicted that miRNA-miRNA interactions of trastuzumab

Figure 5. MMPs related with human breast cancer. Six MMPs of HER2-positive, ESR-positive, triple negative, trastuzumab resistant, tamoxifen resistance and doxorubicin resistance were computed and shown as radar chart.
Figure 6. METS simulation of tamoxifen resistance in human breast cancer. ESR-positive human breast cancer (METS analysis 1) was treated with tamoxifen and then cells obtained a drug resistance as the acquired character (METS analysis 2) (a). From quantum energy level diagram, top level of layers (QCR: 70-80) had compared each other. METS analysis 1. Three miRNAs, miR-34a-5p, let-7a-5p and miR-206 in 70-80 QCR were selected in the simulation of ESR-positive breast cancer (b). Then, target protein genes were presented as the network among miRNAs.

METS analysis 2. Four miRNAs, miR-342-5p, miR-375, miR-320a and miR-378a-5p in 70-80 QCR were analyzed in the next simulation of tamoxifen resistance to be compared with METS analysis 1 (c). Target protein genes were also shown. The clustered genes (red circles) were further characterized by GO cellular component enrichment.
resistance would be implicated in cell membrane signalling from these GOs. Five of the seven miRNAs (AUC; 0.71, p<0.01) have been related with breast cancer [101-103]. Since miR-7-5p and miR-375 are tumour suppressor of breast cancer and both are implicated in trastuzumab resistance of HER2 positive breast cancer, especially, miR-7-5p expression sensitized HER2∆16 to trastuzumab, predicted proteins would be therapeutic targets of HER2-positive breast cancer resistant to trastuzumab [104,105]. In the case of trastuzumab resistance, total target prediction data by METS statistically shows the area under the curve (AUC) 0.90 (P<0.001) in receiver operating characteristic (ROC). Thus, it is suggested that humoral and cellular immune reactions via human monoclonal antibody is controlled by miRNA genes.

Layers of quantum code in miRNAs were investigated in the triple negative breast cancer (Figure 8a). QCR 40-50 in triple negative breast cancer was compared with that in doxorubicin resistant breast cancer. In addition, top layer of doxorubicin was also simulated by METS. Although triple negative breast cancer is characterized by no achievement of complete response with chemotherapy, prediction of chemotherapy response needs to be addressed. In QCR 40-50, upregulation of miR-375-5p would inhibit activation of tumour suppressors, PTEN and a Wnt signalling component, CTNNB1 (Figure 8b). Further, downregulation of miR-200a-5p would increase TP53 suppressor, MDM4. Although amplification of 8p11-12, including LSM1, BAG4 and C8orf4 was observed in approximate 15% of breast cancer, activation of LSM protein component of LSM6, which is involved in mRNA degradation, was also simulated in QCR 40-50. On the other hand, doxorubicin resistant breast cancer upregulated miR-181a-5p and downregulated miR-106a-5p (Figure 8c) [106]. By treatment of doxorubicin, upregulation of miR-181a-5p would have still suppressed expression of anti-apoptotic protein BCL2 and downregulation of miR-106a-5p would have enhanced RB1 tumour suppressor even if in drug resistant cells. Therefore, doxorubicin resistant cancer cells would be likely to result drug sensitive in an anti-cancer state in QCR 40-50; in the top layer of QCR 90-100, however, downregulation of miR-663a would control AP-1 complex in GO cellular component analysis (Figure 8d, red circles genes). In addition, lncRNA H19/miR-675-5p
axis is oncogenic [107]. It is suggested that high layer would administer the low layer, therefore, top layer-targeting agents would be necessary for therapy against doxorubicin resistant breast cancer.

Altogether, drug-resistant phenotypes of breast cancer were classified as 1) the substitution of main players in miRNAs on the same quantum energy level, 2) alteration from the hierarchy longitudinal carcinogenic mechanism, such as ribosomopathy, to the quantum level specific tumorigenic one 3) the higher quantum level of layer would control transform. In the case of statistical analysis in tamoxifen and in doxorubicin, the values of AUC were 0.58 and 0.64, respectively.

**Other chemotherapeutic agents against cancer**

As described above, drug resistance was observed in not only immunotherapy but also chemotherapy against cancers. Cisplatin or carboplatin has been applied for therapy of cancers, such as non-small cell lung cancer, which is approximately 85% of lung cancer, however; drug resistance and decreasing drug sensitivity of chemotherapy exhibit and lead to poor efficacy of it. Fadejeva et al. have reported the summary of implication of miRNA in cisplatin-resistance of non-small cell lung cancer [108]. Cisplatin-resistance was associated with complex mechanisms, apoptosis, drug transport, proliferation, metastasis, DNA damage etc. Thirty-two miRNAs validated were assigned to apoptosis, and 22 to cell proliferation and cell cycle, 7 to DNA damage and 9 to drug transport were reviewed. In the case of oxaliplatin, miR-503-5p was upregulated in the drug resistant colorectal carcinoma [109]. Knockdown of miR-503-5p increased sensitivity to oxaliplatin, therefore miR-503-5p induced oxaliplatin resistance through inhibition of apoptosis by targeting the p53 upregulated modulator protein of apoptosis (PUMA). On the contrary, miR-128 overexpression suppressed paclitaxel-resistant non-small lung cancer stem cells and miR-107 enhanced paclitaxel-sensitivity of non-small lung cancer [110,111]. LncRNA MALAT1 knockdown reversed temozolomide resistance of glioma cells via miR-101 increasing [112]. Together, these data indicated that fail or success of chemotherapy to cancer would depend on alteration of miRNA expression in drug treatment. Thus, METS analysis would be useful to elucidate drug resistance.

**Figure 8.** METS simulation of doxorubicin resistance in human breast cancer. Triple negative human breast cancer (METS analysis 5) was medicated with doxorubicin (a) and drug resistance was emerged (METS analysis 6). The QCR 40-50 was compared with each other. METS analysis 5. The triple negative breast cancer upregulated miR-373-5p and downregulated miR-200a-5p in QCR 40-50 (b). METS analysis 6.1. The network of doxorubicin resistant breast cancer was simulated to be compared with triple negative breast cancer in QCR 40-50 (c). The doxorubicin resistant breast cancer upregulated miR-181a-5p and downregulated miR-106a-5p. METS analysis 6.2. Doxorubicin resistant breast cancer was also simulated in QCR 90-100 (d). Red circled protein gene cluster (enclosed by a large red circle) was analyzed by GO enrichment and it showed as AP-1 complex.
Nutrients

Nutrients of diet could also affect miRNA expression. Quintanilha et al. have summarized the evaluated alteration of miRNAs by nutrients, which are not only polyphenol but also saturated fatty acid (SFA), PUFA, minerals, vitamins, and food-derived miRNAs [113]. Shortly, SFA, palmitic acid increased miR-29a in human myocytes and miR-27b in mouse [114,115], Eicosapentaenoic acid (EPA) and docosahexaenoic acid (DHA) as PUFAs downregulated miR-146a, miR-146b, miR-21, miR-125a, and miR-155 [116]. Food sources of PUFA intervention also altered miRNA profile in the plasma [117]. Omega-3 PUFA, DHA downregulated miR-26a/b in human cholangiocarcinoma cells [118]. MiR-26a/b reduction resulted in inhibit the carcinoma cell proliferation via inhibiting the NAD+ dependent 15-hydroxyprostaglandin dehydrogenase (15-PGDH) through PGE, inactivation, therefore, DHA prevented cholangiocarcinoma growth. In the case of rat model, ω-3 PUFA significantly downregulated miR-19b-3p, miR-146b-5p and miR-183-5p and suppressed inflammation [119]. Zanoaga et al. have summarized breast cancer connection between ω-3 or ω-6 PUFA and miRNA profiles. PUFAs diet could affect miRNA expression, therefore, the environmental factors, such as nutrients have diverse effects against human health as a source of energy as well as genetic information modifier [120]. Vitamin D suppresses miR-155 expression that controls inflammation in patients [121]. Vitamin A (retinoic acid) upregulated miR-10a in breast cancer cells and upregulation of miR-10a suppresses breast cancer cells [122]. Vitamin C modulated mRNA expression of miR-3619-5p, miR-548a-3p, miR-4741, and miR-1825, miR-1208 in human bone marrow stromal cells [123]. Fatty acids and vitamin are nutrients, and some nutrients would bind to RNAs, however, fatty acids are oxidized in mitochondria to release energy and ascorbic acid (vitamin C) is a reducing agent during hydroxylation of collagen. It shows that two nutrients have distinct functions and miRNAs are also multiple functions, which would have the personal diversion, therefore, the relation among nutrients and miRNAs would be the multiple interactions based individual condition on demand. Quite recently, Gong et al. has found the difference expression profiles between American women of African ancestry and of European one, however, they could not determine whether triple negative breast cancer in African ancestry is more aggressive than European one [124]. It means that racing diversion would be linked to miRNA response, in turn, responses of miRNAs to therapeutic agent would also be involved in racing and personal diversion.

Conclusion

Chemotherapeutic and antibody anti-cancer agents, natural substances, nutrients, and miRNA itself in foods affect expression of miRNAs in human. The responses of miRNA to chemicals and antibodies are distinct from in vitro pharmacokinetic reaction; therefore, chemical and antibody substances may have divergent characters in pharmacokinetics with side effect and drug resistance. In general, pharmaceutical company thinks that drugs would be designed not to be affected by personalized variations in pharmacokinetics and the main purpose for their tests would be establish common drug for good sale; however, alteration of miRNA profile means that the environmental substances have an important role for miRNA response to tune our lives and cancer therapy. Since the genetic test with miRNA can be achieved upon cheap and convenient form, a whole series of pharmaceutical drugs including ready commercial ones should be re-validated for personalized medicine by miRNA plus ncRNA expression profiles. Recently, clinical and experimental data have been supplied by open access journals for researchers to investigate miRNA biology, and the big databases of cancer, such as The Cancer Genome Atlas (TCGA) have also been launched into the net-covering the globe. Using these databases, the miRNA target prediction tool has been available for every researcher with equal conditions. We have introduced the quantized miRNA/miRNA interaction and the quantum language of miRNA was participated in the miRNA/target interaction on human cancers. Furthermore, the network of cancer among miRNA/miRNA/target genes, partly lncRNA has clearly been simulated by METS algorithm. On the contrary, drug resistance on approximate 1.4 million women of breast cancer patients is the serious problems in treatment of cancer, but the mechanisms of resistance are acquired, multiple, complex, and not mutually exclusive, and remained to be cleared. To understand the diversity of miRNA response against the chemo-substance, drug resistance in human breast cancer was shown to contextually simulated by METS with quantum multilayer diagram. Trastuzumab resistance simulation resulted significantly high value of AUC. However, neither of tamoxifen and doxorubicin miRNA/miRNA/target context was statistically enough to simulate the drug resistant state of breast cancer because lncRNA and circRNA data were absolute less than microRNA data. Although dynamic computer simulation models of drug resistance in breast cancer showed that in vivo pharmacokinetic effects of drug are much distinct from those of it in vitro, it is simultaneously suggested that miRNA gene expression would be altered by environmental quantum energy or its substances, such as chemicals, antibodies and nutrients. Thus, the METs simulation with quantum language of miRNA would contribute to develop prediction of resistant state, recurrence of cancer and new cancer precious medicine. The quantum language of miRNA against foods and their contents, and drug administration according divergent might be useful for human lives to increase the mortality rate of cancers by inhibition of the incidence on drug resistance, finally by anti-cancer.

Conflicts of interest

The author declares not having conflicts of interest.

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