Combination therapeutics in complex diseases

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Abstract

The biological redundancies in molecular networks of complex diseases limit the efficacy of many single drug therapies. Combination therapeutics, as a common therapeutic method, involve pharmacological intervention using several drugs that interact with multiple targets in the molecular networks of diseases and may achieve better efficacy and/or less toxicity than monotherapy in practice. The development of combination therapeutics is complicated by several critical issues, including identifying multiple targets, targeting strategies and the drug combination. This review summarizes the current achievements in combination therapeutics, with a particular emphasis on the efforts to develop combination therapeutics for complex diseases.

Keywords: combination therapeutics • complex diseases • molecular network

Introduction

Living systems in the human body are interconnected networks of molecular components [1]. Multiple complex molecular networks are involved in the mechanisms of many common diseases, especially in complex diseases (e.g., arthritis, diabetes, heart disease, cancer and asthma) [2–6]. The complexity of these diseases limits the efficacy of commonly used single molecular drug therapeutics, which has led to a critical situation in new drug discovery in recent years [7]. The strategy of combination therapeutics has been revisited with the hope of addressing this issue [8].

The number of studies investigating combination therapeutics is increasing annually (Fig. 1). Moreover, some drug combinations have entered clinical trials for the treatment of complex diseases (Table 1). This review will summarize the current achievements in combination therapeutics in complex diseases, with a particular focus on the computational strategy for drug combination.

Combination therapeutics and drugs with multiple targets

Combination therapeutics are concerted pharmacological interventions consisting of several drugs that interact with multiple targets. The combination of drugs may be antagonistic, additive or synergistic.
if the combined effect is less than, equal to or greater than the sum of
the individual drugs, respectively [9]. Synergistic drug combinations
have numerous advantages over monotherapy, including increased
efficacy, decreased dosage with equal efficacy, reduced side effects
and reduced drug resistance [10]. These advantages are mainly due
to the targeting of multiple molecular networks in the human body.
Complex molecular networks are involved in various biological func-
tions of the human body. For example, bone homeostasis, the pro-
cess of creating new bone and removing old bone [11], has been
found to be regulated by the WNT protein signalling pathway [12], the
Notch signalling pathway [13], the RANK signalling pathway [14] and
the bone morphogenetic protein (BMP) signalling pathway [15].
Moreover, the molecular networks in the human body are also likely
to be modulated by redundant pathways, which can compensate for
one another if either one is inhibited. For example, BMP induces tran-
scriptional activation mediated by SMAD1, SMAD5 and SMAD8,
which functionally compensate for each other in various biological
processes such as bone formation [16], trigeminal ganglia subtype
specification [17] and stem cell differentiation [18]. Such pathways
are therefore optimally inhibited by multiple drugs that block all
SMAD1/5/8 pathways, which ensures that the other pathways do not
partially or completely compensate for the inhibition [16]. Blocking
the upstream signal (for example, the BMP receptor in this case)
would not be appropriate because although it could inhibit all SMAD1/

<table>
<thead>
<tr>
<th>Disease</th>
<th>Combination therapeutics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypertension</td>
<td>Nebivolol + valsartan [33]</td>
</tr>
<tr>
<td>Pyelonephritis</td>
<td>Ceftolozane + tazobactam [98]</td>
</tr>
<tr>
<td>Membranous nephropathy</td>
<td>Prednisolone + chlorambucil [99]</td>
</tr>
<tr>
<td>Diabetes</td>
<td>Dulaglutide + lispro [100]</td>
</tr>
<tr>
<td>Rheumatoid arthritis</td>
<td>Adalimumab + methotrexate [101]; Tofacitinib + methotrexate [102]</td>
</tr>
<tr>
<td>Myeloma</td>
<td>Lenalidomide + dexamethasone + carfilzomib [103]; Lenalidomide + dexamethasone [104, 105]</td>
</tr>
<tr>
<td>Colorectal cancer</td>
<td>Capecitabine + bevacizumab [106]; FOLFOXIRI (fluorouracil, leucovorin, oxaliplatin, and irinotecan)+ bevacizumab [107]</td>
</tr>
<tr>
<td>Pancreatic cancer</td>
<td>Gemcitabine + nanoparticle albumin-bound (nab)-paclitaxel (Abraxane) [108–110]</td>
</tr>
<tr>
<td>Chronic lymphocytic leukaemia</td>
<td>Obinutuzumab + chlorambucil [111]</td>
</tr>
<tr>
<td>B-cell lymphoma</td>
<td>Rituximab + cyclophosphamide + doxorubicin + vincristine + prednisolone [112]</td>
</tr>
<tr>
<td>Breast cancer</td>
<td>Pertuzumab + trastuzumab + docetaxel [113]; Paclitaxel + trastuzumab [114]</td>
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<tr>
<td>Melanoma</td>
<td>Trametinib + dabrafenib [115, 116]; Nivolumab + Ipilimumab [117, 118]; Dabrafenib + trametinib [119]</td>
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Fig. 1 Statistics of combination therapeutic studies by year. The data were obtained from the PubMed database.
In combination therapeutics, one drug might also target multiple functions [19]. A drug that is designed to target multiple processes is an adjusted combination therapeutic in single drug form [20]. Multi-target drugs might overcome molecular heterogeneity within or between patients diagnosed with complex diseases, such as non-small-cell lung cancer [21], gastrointestinal stromal tumours [22] and pulmonary fibrosis [23], and would therefore have a better chance of successful clinical development, especially when biomarkers for patient selection are uncertain. However, such a multi-target drug might not be ideal for reasonable therapeutic strategies based on specific molecular profiles of diseases. Because these drugs are not well ‘designed’ to interact with a particular set of targets, they may not achieve optimal inhibition of the specific individual targets in a particular disease [24]. For example, during kinase inhibition, screening to detect the inhibitory activities of particular kinases of interest resulted in the identification of multi-targeted kinase inhibitors. However, additional kinases with structural similarities will be inhibited, potentially leading to additional toxicity because cumulative target and off-target inhibition has a broader, and perhaps less predictable, effect on cellular functions [24, 25]. For example, Sorafenib, a multi-target tyrosine kinase inhibitor with potent anti-angiogenic activity that is used to treat solid tumours, induces irreversible pancreatic atrophy in patients after long-term administration [26].

In contrast, the combination of more specific targeted drugs may be more suitable for therapeutics tailored to individual patients based on the specific molecular disease profiles [27]. Physicians might adjust the concentrations of drugs within a combination therapeutic to enhance targeted inhibition and to optimize synergy between drugs [28]. The toxicity of the combination might also be more predictable due to limited off-target effects [29].

Drug selection for combination therapeutics

Currently, most of the drugs in combination therapeutics are chemosynthetic molecules. For example, Ser-Ap-Es, which has been used for the treatment of hypertension since the 1950s, is a triple-chemical-drug combination that includes reserpine, apresoline and hydrochlorothiazide. Hypertension is a complex cardiovascular disease, and approximately 30% of the adult population in the United States suffers from this condition [30]. Law et al. have suggested that low-dose combinations from the five major classes of antihypertensive drugs (diuretics, beta-blockers, angiotensin-receptor blockers, calcium-channel blockers and angiotensin-converting enzyme inhibitors) are more effective for reducing blood pressure and associated with fewer side effects than monotherapy [31]. Taylor and Ragbir evaluated some triple-drug combinations and showed that they were well-tolerated with a low incidence of side effects [32]. Recently, a fixed-dose combination of a vasodilating β-blocker (nebivolol) and an angiotensin II receptor blocker (valsartan) was shown to be an effective and well-tolerated treatment option for patients with hypertension in an 8-week, phase 3, multi-centre, randomized, double-blind, placebo-controlled, parallel-group trial [33].

Bioengineering products are becoming another important source of drugs for combination therapeutics. Monoclonal antibodies are the most common form of this type of drug [34]. For example, trastuzumab, a monoclonal antibody that interferes with the HER2 receptor, has been used in combination with cisplatin to inhibit the progression of gastric cancer [35]. Cisplatin induces DNA damage and apoptosis, which may be attenuated by DNA repair systems [36]. Trastuzumab suppresses the DNA repair pathway and the PI3K-AKT pathway [37] to increase cellular apoptosis [38]. The combination of ramucirumab, a human monoclonal antibody against VEGFR2, and paclitaxel significantly increased the overall survival of gastric cancer patients in a randomized, placebo-controlled, double-blind, phase 3 trial in 27 countries in North and South America, Europe, Asia and Australia [39].

Herbs have been widely used in eastern Asia for thousands of years. Herbal compounds would be good candidates for combination therapeutics. There are already several successful cases examining this theory. Paclitaxel, a drug developed from a compound isolated from the bark of the pacific yew tree, Taxus brevifolia, is now widely used in combination therapeutics for cancer treatment. The combination of paclitaxel and bortezomib, a proteasome inhibitor that modulates apoptosis and the cell cycle by disrupting protein degradation, improves overall survival of non-small cell lung cancer patients [40]. Total glucosides of peony, active compounds extracted from the roots of the herb Paeonia lactiflora Pall, reduce hepatotoxicity in combination with leflunomide and methotrexate in patients with active rheumatoid arthritis [41]. In China, the herb Artemisia annua is used to treat fever and malaria. In 1971, artemisinin was isolated from the leafy portions of the A. annua plant and found to be effective for the treatment of malaria [42]. Artemisinin-combination therapy is now recommended as a first-line treatment for falciparum malaria worldwide, and fixed-dose combinations are preferred by the WHO. In addition, some combinations of herbal compounds are in preclinical phases. Deng et al. have indicated that the combination of salvianolic acid B and ginsenoside Rg1, which are derived from roots of Salvia miltiorrhiza and Panax notoginseng, respectively, improve the viability of cardiac myocytes in rats with ischaemia/reperfusion injury [43]. Zhang et al. have found that tetramethylpyrazine, a compound isolated from Ligusticum wallichii Franchat, ameliorates oxidative organ injury associated with methotrexate treatment in patients with rheumatoid arthritis [44]. These examples illustrate the possibility of utilizing herbal compounds in combination therapeutics. Therefore, herb might be a rich resource for the discovery of combination therapeutics.

Drug combinations are much more than synergistic

In the clinic, physicians often face the following pragmatic question: should I use a second drug as an add-on instead of using more of the first drug? In the case of complex diseases, the answer will often be that an increased amount of the first drug cannot produce the effect that is desired by the physician, but this effect can be achieved with
combination treatment [28]. Moreover, in some cases, the addition of more of the same drug will induce unacceptable negative influence. For example, when the dosage of the first drug is slightly under the threshold of toxicity, even a small dosage increase may be unacceptable [45]. Therefore, combining the first mediation with a second drug that produces non-overlapping toxicities may be beneficial, even when the combined effects on efficacy are only Loewe additive [34, 35], which refers to a model of drug combination and will be introduced in the next paragraph. Ultimately, the ideal design of combination therapeutics must reflect the requirement of the physician that, when drugs are used in combination, an extra benefit is observed that cannot be achieved by using the constituent drugs alone. The extra benefit is mainly derived from the synergistic effects of the combined drugs (Fig. 2A).

The word synergy is derived from the Greek words sunergia, which means ‘cooperation’, and sunergos, which means ‘working together’ [19]. Modern pharmacology has embraced this concept to emphasize the idea that a synergistic combination provides greater effects than those predicted by simply adding the effects of individual parts [10]. Numerous investigators have attempted to provide a framework to calculate the additive effects among constituents [46–48]. Among these works, the Loewe additivity [49, 50] and Bliss independence [46, 51] models are two of the most useful frameworks of synergy.

The Loewe additivity model assumes that two drugs act through similar mechanisms. Equitotent dose ratios determine the effects of each drug and their combination. According to this model, the combined activity level, $L_{x+y}$, of the two drugs $X$ and $Y$ at the concentrations, $C_x$ and $C_y$, satisfies the equation, \[
\frac{C_x}{C_0} + \frac{C_y}{C_0} = 1.
\]
In this equation, $C_x$ and $C_y$ are the single-agent concentrations for $X$ and $Y$, which individually produce an activity of $L_x$ and $L_y$, respectively.

The Bliss independence model assumes that two drugs act through independent mechanisms. The combination effect in this model is represented as the union of two probabilistically independent events [46]. According to this model, the combined activity, $L_{x+y}$, of two compounds $X$ and $Y$ at the concentrations, $C_x$ and $C_y$, satisfies the equation, $L_{x+y} = L_x + L_y - L_xL_y$. In this equation, $L_x$ and $L_y$ are the single-agent activity levels at concentrations $C_x$ and $C_y$.

These two models produce different outcomes, and only the Loewe additivity model correctly predicts the cases in which the two compounds are actually the same constituent. However, the debate continues over which model performs better in cases involving noisy clinical data and uncertain therapeutic mechanisms [46, 52, 53].

Nonetheless, synergy is not the only reason for drug combinations. Sometimes the combination of antagonistic drugs may also be beneficial (Fig. 2B). In fact, the combination can have two conflicting effects: it may reduce the evolution of drug resistance because it cures the disease faster, thereby limiting the time window available for drug-resistant mutations to accumulate, but it may also increase the selective advantage of drug-resistant mutants [54]. If a strong resource competition is present, the latter effect will dominate [54]. In fact, under these conditions, antagonistic drug combinations, which are less potent than the sum of the components, have been shown to limit the evolution of drug resistance [55–57]. The advantage of antagonistic combinations over synergistic combinations is the result of a reduced fitness gain of drug-resistant mutants (Fig. 2B) [55–57].

In addition to synergistic and antagonistic drug interactions, collateral responses among drugs also play a role in combination therapeutics (Fig. 2C) [58]. Collateral response (sensitivity or resistance), also known as cross-resistance, is the ability of one drug to increase or decrease sensitivity to another drug (Fig. 2C). The study of collateral susceptibility changes started in the early 1950s [59] and has been reported for many different drugs in subsequent years [60–62]. Recently, Munck et al. studied the development of resistance against drug combinations by investigating collateral responses to component drugs [58]. The authors provided a framework for the rational selection of drug combinations that limit resistance evolution using Escherichia coli.

In general, the benefits of combination therapeutics mainly result from synergistic effects. Antagonistic effects and collateral responses may also produce extra benefits by decreasing drug resistance. Therefore, combination therapeutics can be more complicated than simply taking different drugs together. To investigate the rationale of combination therapeutics, preclinical combinational studies should be performed prior to the initiation of clinical trials.

**The development of combination therapeutics in cell lines and animal models**

A common strategy to investigate combination therapeutics prior to the initiation of clinical trials is through the use of cell lines and animal models. These models provide important insights into the mechanisms of action and interactions between drugs, which are important in the preparation for clinical trials [63]. For example, DiCosimo et al. estimated the efficacy of combined ridaforolimus and dalotuzumab for the treatment of breast cancer in human cell lines and mouse models, and the results of those studies led to the initiation of a phase I clinical trial [64]. In fact, preclinical studies make up the majority of studies investigating (Fig. 1) and play an essential role in the development of combination therapeutics [65]. In general, preclinical studies of combination therapeutics rapidly assess the synergy of combined drugs and explore physical interactions among them. They also evaluate anti-disease activity, mechanisms of action and additional factors, such as the effects of the drugs on their targets and their pharmacokinetics [66].
However, despite the large number of preclinical combination therapeutics studies, the number of successful clinical trials is limited (Fig. 1). It appears that the success or failure of clinical trials examining combination therapeutics can hardly be predicted on the basis of published preclinical data [67]. The lack of concordance between preclinical studies and clinical trials is not surprising. Preclinical studies are limited in experimental design and animal models [63]. In experiments that are designed to evaluate the activity of combined drugs, assessments are performed with limited consideration of how the drugs are likely to function in patients [83]. Such evaluations can provide insights into the mechanisms responsible for anti-disease actions and the interactions between the drugs within the parameters of the experiment. However, interpretation of the evaluations and the subsequent design of clinical trials for complex diseases must encompass numerous variables that might exceed the designed parameters of preclinical experiments, such as drug dose, scheduling and route of administration [68].

Even when researchers have thoroughly designed all of the parameters of preclinical experiments for combination therapeutics, the inherent limitations of animal models continue to weaken the correlation between laboratory and clinical outcome. The most important limitation is that the drug targets are host components and are therefore of animal, not human, origin [65]. In other words, a drug that works well in an animal model may not necessarily work in humans. After assessing the correlation between cancer animal models and the clinical activity of single cytotoxic drugs, previous studies have suggested that the predictive value of cytotoxic drugs in murine models is limited [69, 70]. Limiting experimentation to a few animal models cannot reflect the diversity of complex human diseases and may explain the failure of clinical trials based on preclinical data [67]. A detailed understanding of human biology may enable the design of a priori combination therapeutics to increase the success rate of clinical trials. A growing number of scientists in the field of therapeutic discovery are attempting to understand biology and disease using computational approaches [71].

The development of combination therapeutics using computational approaches

Computational approaches, particularly those integrating data analysis and artificial intelligence, may assist in the design of combination therapeutics by generating drug–target interaction networks and predicting drug combinations. Computation assistant construction of drug–target interaction networks mainly refer to computational algorithms and programs to reveal molecular mechanisms and connections between drugs and targets in a dynamic network [72]. These methods can be used to explore the therapeutic mechanisms underlying drug treatment at the molecular level [73]. Common methods of this type include knowledge- and ligand-based approaches.

Knowledge-based approaches are high-throughput computational methods of artificial intelligence that enable computers to learn from the available knowledge, including drug chemistry and structural information, as well as drug–target networks, to predict new knowledge, such as new drug indications, targets and drug–target interactions [74]. These approaches can be classified into unsupervised and supervised methods [75]. The unsupervised methods extract and predict patterns and interactions based on a series of input variables. The commonly used unsupervised methods include clustering, data compression and outlier detection methods such as principal-component-based methods [76]. For example, Guimera et al. studied novel drug–drug interactions using a large-scale unsupervised method that deals with various types of data to accurately predict adverse, synergistic and antagonistic drug interactions [77]. Kissa et al. examined drug–gene associations and assisted in drug repositioning with a fully corpus-based unsupervised method that utilizes the available knowledge [78]. The supervised methods divide the data into training and validation datasets, and they finalize a robust model that can be used to predict the binding probability between drugs and targets [79]. The most widely used supervised methods are the Bayesian model [80], the support vector machine [81] and the decision tree [82]. These methods can be combined if appropriate. For example, Kim et al. predicted drug–target interactions by applying two machine learning approaches, a support vector machine and a kernel-based L1-norm regularized logistic regression, to evaluate drug–drug interaction data [83].

Ligands are small molecules that bind to a site on a target protein to alter the biological functions of the protein. Ligand-based approaches predict drug–target interactions by means of chemical or structural ligand similarities, with the basic assumption that similar ligands tend to bind to similar targets [84]. Ligand-based approaches are highly dependent on the availability of chemical and structural information on targets, from either wet experiments or numerical simulations [85, 86]. The similarity ensemble approach [87], the pharmacophore model [88], docking [89] and quantitative structure-activity relationships [90] are commonly used methods. Based on these types of methods, Yco et al. identified sepiapterin reductase as a new therapeutic target for the treatment of human neuroblastoma [91].

Knowledge- and ligand-based approaches are often combined to achieve better results. Afzel et al. combined multiple methods to construct drug–target predictions and obtained more applicable results [92]. These computational approaches accelerate the acquisition of drug–target data. The accumulation of these data, once again, promotes novel predictions of drugs targets [93].

In addition to accelerating the accumulation of drug targets, computational approaches directly predict new combination therapeutics. Bansal et al. assessed 32 computational approaches and found that the computational prediction of drug combination activity is possible [94]. The prediction of new combination therapeutics should include network pharmacology that encompasses systems biology, network analysis, connectivity, redundancy and pleiotropy [95]. Gu et al. predicted a drug combination in the context of the pathway network of the biological process [96]. The computational approach may become an important complement to traditional preclinical studies. It is possible that computational approaches will be used to identify promising combinations that can be tested in vitro to further determine the combinations, which would be advantageous compared with random...
Combination screening in vitro. This method may even replace in vitro screening in cell lines in the future, but in vivo efficacy testing would be still required. Large and complex biological data and multiple computational methods are often used in studies focused on the prediction of combination therapeutics [97]. With the accumulation of chemical, biological, ‘omics’ data and the advancement of artificial intelligence, computational approaches will acquire a more prominent role in the development of combination therapeutics for complex diseases.

Conclusions

The use of combination therapeutics is an appropriate approach to address the complexity of human diseases. The advantages of combination therapeutics are mainly due to the ability of combination products to act on multiple targets. Chemical drugs, biological drugs and drugs from natural products are widely used in combination therapeutics. Drug interactions, including synergistic and antagonistic drug interactions, as well as collateral responses, might influence the therapeutic actions of drug combinations. Preclinical studies in cell and animal models have been performed to evaluate the effects of drug combinations. However, subsequent successful clinical trials of drug combinations are limited, potentially because the limited experimental conditions in preclinical studies cannot reflect the diversity of complex diseases. Computational approaches can complement traditional preclinical studies. The use of computational approaches not only facilitates an understanding of drug interactions, but it also promotes the discovery of new combination therapeutics. Computational approaches might help to define the number of candidate combinations for further in vitro and in vivo investigations. In future, the accumulation of multiple types of human ‘omics’ data, including genomics, transcriptomics, proteomics and metabolome, among others, will deepen our understanding of complex diseases and promote the computational discovery of combination therapeutics.

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Conflict of interest

The authors declare that there are no conflicts of interest.

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