Repurposing approved drugs on the pathway to novel therapies

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Abstract
The time and cost of developing new drugs have led many groups to limit their search for therapeutics to compounds that have previously been approved for human use. Many “repurposed” drugs, such as derivatives of thalidomide, antibiotics, and antivirals have had clinical success in treatment areas well beyond their original approved use. These include applications in treating antibiotic-resistant organisms, viruses, cancers and to prevent burn scarring. The major theoretical justification for reusing approved drugs is that they have known modes of action and controllable side effects. Coadministering antibiotics with inhibitors of bacterial toxins or enzymes that mediate multidrug resistance can greatly enhance their activity. Drugs that control host cell pathways, including inflammation, tumor necrosis factor, interferons, and autophagy, can reduce the “cytokine storm” response to injury, control infection, and aid in cancer therapy. An active compound, even if previously approved for human use, will be a poor clinical candidate if it lacks specificity for the new target, has poor solubility or can cause serious side effects. Synergistic combinations can reduce the dosages of the individual components to lower reactivity. Preclinical analysis should take into account that severely ill patients with comorbidities will be more sensitive to side effects than healthy trial subjects. Once an active, approved drug has been identified, collaboration with medicinal chemists can aid in finding derivatives with better physicochemical properties, specificity, and efficacy, to provide novel therapies for cancers, emerging and rare diseases.
INTRODUCTION

The earliest drugs were derived from natural products or compounds that could easily be made chemically. Many of these molecules are still being used today, including aspirin, steroids, and antibiotics produced by microorganisms. In light of the problems with bringing a new drug to market, many researchers consider screening Food and Drug Administration (FDA)-approved drugs for repurposing a lower risk than de novo drug design. Searching for specific molecules for a novel drug target may be perceived as unrealistically slow, especially during emerging disease outbreaks. Thus, the approximately 4000 compounds used in the clinic are now being reinvestigated, to see what other treatments they could provide for difficult to treat conditions. Searching approved drugs for new purposes has been aided by commercially available compilations of drugs approved by the FDA, such as the Prestwick Chemical Library, the Library of Pharmacologically Active Compounds (LOPAC1280) and others, or specific libraries assembled for various purposes, such as those targeting RNA-protein interactions or kinases. The “Drug Repurposing Hub” (http://www.broadinstitute.org/repurposing), consisting of both a reliable physical source for approved drugs, as well as a virtual library describing their characteristics, will certainly help to identify new uses for the existing pantheon. There are also a variety of other publicly available resources that can assist repurposing, including RepurposingDB, Re:fine Drugs, or repoDB.

The most common reason for repurposing is that approved drugs have already been tested for safety in humans, meaning they should enter the clinic quickly. As discussed in Section 2.1, many older drugs, even those with known serious side effects, have indeed been reintroduced to the clinic for purposes beyond their original indications. The section highlights how, despite a disastrous first introduction and withdrawal from the clinic in the last century, thalidomide and its derivatives are now used to treat a variety of serious health threats. While drugs designed to treat many indications have been introduced into cancer therapy, the thalidomide story illustrates how even a drug with serious known side effects can be successful. It is also an example of how simple changes in structure can generate a much more active molecule, showing how repurposing can aid in the design of novel drugs (section 3). Many other examples are given of drugs that have found new uses, far from their initial indications.

Rapid application of effective antibiotics for bacterial infections can greatly decrease the length of hospitalization and risk of death. The rise of multidrug-resistant bacteria, as well as the paucity of novel antibiotics, has led to many efforts to make the current arsenal more potent by combining older antibiotics. Section 2.2 highlights these strategies, as well as efforts to combine antibiotics with agents that target bacterial toxins or block bacterial enzymes from exporting or degrading antibiotics. In some cases, simply changing the mode of application of a drug is sufficient to enhance activity.

Another impetus for repurposing is to find broad-spectrum treatments for emerging infections by targeting host cell pathways rather than the infectious agent directly. Ways to complement antibiotics and antivirals with drugs that enhance host cell resistance mechanisms have thus become an active and promising area for study, discussed in Section 2.3. One driving force for this is the ability of pathogens to quickly develop resistance to targeted inhibitors. For example, drug resistance has developed to two of four FDA-approved antivirals against influenza. Activating the host immune system is especially important in looking for new ways to treat chronic infections such as tuberculosis. New ways must also be found for treating viruses, including Ebola, that have developed mechanisms to evade host immune systems. Paradoxically, the repurposing wave is occurring just as the clinical success of direct-acting antivirals (DAAs) against Hepatitis C virus has shown the power of high throughput screening coupled with structure-based, rational design.
Section 3 is devoted to the difficulties in repurposing drugs and natural products, and how older drugs may be repurposed to overcome these. Many approved drugs have low solubility, lack specificity, and may require unrealistic dosing for uses beyond their initial target (Section 3.1). Dosages that may be well tolerated in trials conducted using healthy subjects (or in the case of animal studies, healthy until infected) may not be achievable in the clinic. Patients in the clinic or field hospital are very sick when they present and have comorbidities. Burn, cancer, Ebola, and other emerging infection victims are fragile patients. They will thus be more likely to suffer side effects at high doses. Collaboration with medicinal chemists at an early stage in development can insure more specific and potent therapies for testing. In Section 3.2, a few examples show how derivatives of older drugs, with better pharmaceutical properties and selectivity, can bring new treatments into the clinic. These derivatives can add to the continuously expanding drug pantheon.

2 FINDING NEW USES FOR APPROVED DRUGS

2.1 Drugs with clinical success for purposes beyond their initial approved use

2.1.1 The many current uses of thalidomide

The many current uses of thalidomide represent perhaps the most spectacular example of how an old drug can assume new roles.22 Thalidomide (Table 1) was originally introduced by Grünenthal Chemie in the mid-1950s, and by 1957 was available over the counter in several European countries for treating insomnia and morning sickness. The drug was not approved by the US-FDA, as the young examiner assigned to the thalidomide application noted defects in its safety profile. She was especially concerned about severe peripheral neuritis reported as a possible side effect.23 Thus, even without teratogenicity testing, thalidomide’s potential to cause dangerous side effects was clear before 1960. Thalidomide was withdrawn worldwide in 1962 only after it became notorious for causing severe birth defects. The drug’s other associated side effects, including rash, tremor, hypothyroidism, hypocalcemia, hyperkalemia, and toxic epidermal necrolysis (in 3%-4% of patients) would seem to have doomed it to a mere cautionary tale in the history of pharmacy.

Yet thalidomide soon rose from the ashes. By the mid-1960s, it was being used to treat skin lesions and granulomas associated with leprosy. Today, thalidomide is part of the first-line treatment for Hanson’s disease. Curiously, its ability to bind to cereblon,24 a protein involved in limb outgrowth, which is believed to be at the root of its teratogenic effects, may also enable it to control the growth of multiple myeloma cells.25 More potent derivatives, such as lenalidomide and pomalidomide (Table 1) have been approved by the FDA to treat multiple myeloma,25 mantle cell lymphoma, and myelodysplastic syndromes associated with the deletion 5q abnormality.

Thalidomide and its derivatives inhibit secretion of tumor necrosis factor (TNF)-α and other inflammatory cytokines,26 by direct and indirect effects on cells.27 This has led to many tests for their abilities to ease inflammation. Low dose thalidomide (50-150 mg/day) alleviated the symptoms of patients with Crohn’s disease who were refractory to many other treatments.28 Thalidomide also alleviates the cytokine storm induced by bacterial infections.29 Animal studies indicate thio-derivatives, such as 3.6'-dithiothalidomide, might be useful for the treatment of aneurysms30 and traumatic brain injury.31 The more specific PDE4 inhibitor, apremilast,32,33 can alleviate several diseases associated with overexpression of TNF-α, such as psoriasis, lupus erythematosus, and rheumatoid arthritis.34

Lenalidomide treatment may even restore color to gray hair.35 While this might be related to other effects of TNF-α on regulating hair growth,36,37 it should be noted that other therapies, such as tyrosine kinase inhibitors,38 may also affect hair color. “Repigmentation”, in this case, darkening of hair, has also been seen during therapy with PD-L1 inhibitors. The authors describe repigmentation as a side effect and possible clinical marker for successful treatment.39
2.1.2 | Many other older drugs are being repurposed

Recent government support has led to many more approved drugs being tested for applications far beyond their initial use. For example, drugs that have “antitussive, sedative, analgesic, antipyretic, antiarthritic, anesthetic, antidiabetic, muscle relaxant, immunosuppressant, antibiotic, antiepileptic, cardioprotective, antihypertensive, erectile function enhancing, or angina relieving” properties may be used as adjuvant therapies in cancer. A few examples of repurposed drugs that are being tested in select groups of patients or have entered clinical trials are shown in Table 2.

The effort to reuse existing drugs for novel therapies is especially important for rare diseases, where the patient population is too small to justify the major effort needed to introduce a new drug. Alternatively, it may be the only way to identify treatments for severe diseases with idiopathic etiology, such as neurofibromatosis.

As Table 2 indicates, repurposing may provide alternative treatments for psychiatric disorders. The antioxidant ebselen has been tested to treat bipolar disorder in patients who cannot tolerate lithium treatment, while loxapine may treat irritability associated with autism. Antiinflammatory drugs, as well as a variety of other repurposed compounds, have also been suggested as alternative treatments for schizophrenic patients. These include acetylsalicylic acid, celecoxib, minocycline, N-acetylcysteine, and nutraceuticals.

### Table 1: Thalidomide and derivatives

| Compound                     | Structure | Human dose     |
|------------------------------|-----------|----------------|
| Thalidomide                  | ![Thalidomide](image) | 50-150 mg/d    |
| Lenalidomide (revlimid)      | ![Lenalidomide](image) | 5-25 mg/d      |
| Pomalidomide (pomalyst)      | ![Pomalidomide](image) | 2-4 mg/d       |
| 3,6′-Dithiothalidomide       | ![3,6′-Dithiothalidomide](image) | Not determined |
| Apremilast (otezia)          | ![Apremilast](image) | Up to 60 mg/d  |

*Note: Compound drawings are from the ZINC database or relevant publications.*

2.2 | Making the current antibiotic arsenal more effective

#### 2.2.1 | Reusing older antibiotics

Natural microbial communities consist of complex mixtures of bacteria, viruses, and fungi, with each organism fighting against the others for dominance. An organism survives by secreting antibiotics and toxins, and by evolving internal enzymes to deactivate compounds that succeed in penetrating its own cell wall or envelope. The most
| Compound                     | Initial use                       | Repurposed for:                          | Mechanism                                                                                                                                                                                                 | Clinical stage/comments                  |
|------------------------------|-----------------------------------|------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------|
| Atorvastatin (generic Lipitor) | Hypercholesterolaemia             | Cavernous angioma                        | Blunts lesion development and hemorrhage through inhibiting RhoA kinase (ROCK)                                                                                                                             | Exploratory; proof of concept\(^{40}\)   |
| Statins                      | Hypercholesterolaemia             | Oncology                                 | Inhibits production of mevalonate and isoprenoids involved in Ras and other small GTPase oncogenes pathways                                                                                               | Clinical failures may be due to bad clinical trial design\(^{31}\) |
| Losartan                     | Blood pressure reduction          | Alzheimer disease                        | Angiotensin 1 receptor antagonist; high blood pressure may exacerbate AD                                                                                                                                 | Preclinical trials\(^{42}\)              |
| Fenofibrate                  | Reduces, triglyceride-rich particles (LDL) in plasma | Reduces macrophage recruitment in abdominal aortic aneurysm | Activates lipoprotein lipase, to remove LDL; reduces the proinflammatory protein osteopontin                                                                                                                | Ongoing randomized controlled trials\(^{43}\) |
| Telmisartan                  | Blood pressure reduction          | Abdominal aortic aneurysm                | PPAR-γ agonist, reduces TGF-β, MMP-9 and other biomarkers associated with progression                                                                                                                        | Ongoing randomized controlled trials\(^{43}\) |
| Amiloride                    | Acid-sensing ion channel antagonist | Secondary progressive multiple sclerosis (SPMS) | These three drugs were selected from seven candidates for clinical testing for repurposing as neuroprotective therapies in MS                                                                                   | Multiple Sclerosis-Secondary Progressive Multi-Arm Randomisation Trial (MS-SMART): a multiarm, phase IIb randomized, double-blind, placebo-controlled clinical trial\(^{44}\) |
| Fluoxetine                   | Serotonin selective reuptake inhibitor (SSRI) |                                           |                                                                                                                                                                                                         | Possible candidate for clinical testing\(^{45}\) |
| Riluzol                      | Glutamate antagonist              |                                           |                                                                                                                                                                                                         | Possible candidate for clinical testing\(^{45}\) |
| Edaravone                    | Neuroprotective agent in acute ischemic stroke and ALS | Multiple sclerosis | Aids in remyelination and neuroprotection                                                                                                                                                                   | Possible candidate for clinical testing\(^{45}\) |
| Dexamethasone                | ALS and other neurological diseases; phase 3 trials did not meet endpoint | Hypereosinophilic syndromes              | Serendipitous observation that patients in the ALS studies had reduced eosinophils.                                                                                                                                 | 12 Wk/10 patient phase II trial showed glucocorticoid doses, paving the way for phase III\(^{46}\); NCT02101138 |
| Disulfiram (antabuse)        | Reduces ethanol tolerance in alcoholism | Metastatic breast cancer & Alzheimer disease | Blocks acetaldehyde dehydrogenase; modulates ADAM10, an α-secretase in neurons which cleaves amyloid precursor protein to sAPP-α                                                                                                                                 | In testing, with copper, for breast cancer: NCT03323346 basis for testing in AD\(^{47}\) |

(Continues)
| Compound | Initial use | Repurposed for: | Mechanism | Clinical stage/comments |
|----------|-------------|-----------------|-----------|-------------------------|
| Metformin | Diabetes | Anti-nonsmall cell lung cancer, Augmented resistance in aging | Decreases hepatic glucose production and enhances insulin sensitivity. The effects in other diseases were observed in many users | Phase II for NSCL, trial; safe but not effective. However, only 14 of 50 patients were enrolled resistance in aging trial (MASTER) |
| γ-Secretase inhibitors (GSI) | Alzheimer disease: prevent amyloid precursor cleavage | Several inhibitors are being testing against a variety of cancers | GSIs can inhibit NOTCH1 signaling by inhibiting the secretase, some also inhibit signal peptide peptidases | Ongoing clinical trials (NCT01981551, NCT03785964, NCT03691207), as well as in combination with other cancer drugs or Car T-cell therapy (NCT 03502577) |
| Saracatinib (AZD-0530) | Cancer therapy | Mild to moderate Alzheimer disease | Inhibits SRC, Bcr-Abl and Fyn Kinase, the latter may contribute to synaptopo-toxicity in AD | Phase Ib-II trials for AD indicated the drug was safe but efficacy was unclear |
| Mibefradil (Posicor) | Antihypertensive, calcium channel blocker | Short term use as an adjuvant in cancer therapy | Enhances action of anticancer agents and radiation, tested especially for glioblastoma | Several trials for use in combination therapies |
| Nelfinavir | HIV protease inhibitor | Solid tumors | Inhibits endogenous Akt activity in cancer cells | Phase | NCT01445106 |
| Human Albumin | Blood additive | Immunorestitution | Binds and inactivates prostaglandins | ATTIRE study ongoing in England to restore serum albumin to >30 g/L by concentrated infusions |
| Ebselen | Antioxidant, (mimics glutathione peroxidase) | Bipolar disorder | Ebselen and lithium both inhibit IMPase and reduce glutamate and inositol levels in brain areas | Tests in healthy volunteers using up to 3600 mg dose |
| Loxapine | Antipsychotic and antischizophrenia | Irritability associated with autism | Rebalances dopamine and serotonin levels; increase brain-derived neurotrophic factor (BDNF) | 12 Wk open trial |
| Nitazoxanide | Antiprotozoal agent | Influenza | First-in-class broad-spectrum thiazolide antiviral inhibits strains resistant to neuraminidase inhibitors | Phase III: NCT03336619 |

Note: Numbers beginning with NCT are ClinicalTrials.gov ID’s for related studies. See for antibiotics and other drugs that have been repurposed for cancer therapy. Abbreviation: TNF, tumor necrosis factor.
Clinically important microbes are survivors of these battles, meaning they typically possess powerful defensive weapons consisting of an array of toxins, enzymes to degrade or export antibiotics, and other tools to deal with serum proteases and the mammalian immune system. In the face of antibiotic treatment, the enzymes in this arsenal evolve to meet the new challenge.

In the past, older antibiotics were constantly being superseded by newer ones, designed to evade the latest resistance mechanism found in clinical isolates. However, many companies have dropped programs to develop new antiinfectives, due to the time and cost of clinical trials. The lack of new drugs has given a strong incentive to repurpose older antibiotics for multidrug-resistant strains or to combine them with agents that can target several different pathways in the pathogen, or activate host cell resistance pathways. This should lead to enhanced activity and lower the chances for development of resistance.60

Reverting to older antibiotics can, in some cases, overcome antibiotic resistance. For example, TB strains resistant to currently used drug combinations are found in all parts of the world. One of the most useful TB drugs, isoniazid, inhibits the enzyme InhA, blocking the synthesis of the mycolic acids that make up the cell wall of the mycobacterium. Much of the resistance to isoniazid, a prodrug, comes from mutations in the bacterial enzymes required to activate it. It has recently been found that an older natural product antibiotic, pyridomycin, discovered in the 1950s, inhibits InhA directly. Thus it can be used to treat infections with resistant TB strains that survive by their lack of ability to activate isoniazid.61 Understanding this mechanism of action may also advance the use of pyridomycin in conjunction with other drugs.62

2.2.2 Antibiotic adjuvants and toxin inhibitors

The era of antibiotic resistance has also brought a new interest in combining older antibiotics with compounds that can enhance their activity.63 Many compounds have been developed to specifically block bacterial enzymes that mediate resistance.64 Combining penicillin derivatives with β-lactamase inhibitors can overcome bacterial resistance in many cases.19 For example, one can pair piperacillin with tazobactam, a competitive inhibitor of β-lactamases that protects the paired antibiotic from degradation. The mechanisms of β-lactamases and their respective inhibitors that have been successfully combined with antibiotics are discussed in several comprehensive reviews.65,66

While combination therapy has advantages, the rapid rise of resistance mutations in bacteria67 can complicate the choice of combinations to yield the best synergy among treatments.68 Determining the most effective adjuvants to complement antibiotics to better treat resistant bacteria should be aided by a new “antibiotic resistance platform” (ARP). The ARP is an array of transformed Escherichia coli that expresses one of greater than 40 different known antibiotic resistance genes. This allows screening for compounds that either target the resistance gene or otherwise potentiate the antibacterial activity of a given antibiotic.69 Other efforts are being made to design combination therapies for fungal infections.70

Cotreatments with toxin inhibitors

Combining antibiotics with agents that inhibit toxins or other effectors that contribute to their pathogenicity can also enhance their activity. Tissue damage and immune activation are caused by bacterially produced toxins, including proteases, pyrolytic toxins, “superantigens”71 and those that elevate tissue cyclic adenosine monophosphate levels.72 Antibiotics can have additional activities that may lead to synergy or antagonism of toxin production. For example, β-lactam antibiotics can increase the production of bacterial toxins in Staphylococcus aureus, while those that inhibit protein synthesis, such as clindamycin, greatly reduce them.73 As toxins mediate pathogenic effects even after the death of the bacteria, direct inhibitors of bacterial toxins may aid in controlling symptoms when used alone or in combination with antibiotics.15,74-76 Further, as discussed in the next sections, patients may benefit from alternative application modes and cotreatment with antibiotics and other compounds designed to directly target pathways in human cells.
2.2.3 | Changing the mode of application

The application mode of antibiotics can also affect their activity. While oral availability is prized, inpatients are frequently treated with antibiotic infusions or direct injection.

Inhaling a drug may give completely different activities than injection or infusion. Recent experiments in pigs with deleted cystic fibrosis transmembrane receptor (CFTR) have shown that an aerosol treatment with amphotericin B, a toxic antifungal agent, formulated in liposomes with cholesterol, restored airway pH levels. The authors suggest that similar aerosol treatment would help CF patients with many different CFTR mutations.77 Similarly, inhalation may also improve the ability of type-1 interferon to treat Epstein-Barr virus.78 interferon (IFN)-γ, which has mechanisms of action that differ from type-1 interferons,79 such as in its ability to directly control ribonucleases,80,81 was originally used as an antiviral and immune activator. Currently, it is used via subcutaneous injection to treat chronic granulomatous disease and osteopetrosis. In a controlled clinical trial, inhaled IFN-γ could control TB, while parenteral IFN-γ did not. These results indicate that macrophages can be effectively immune-stimulated by aerosol therapy. Treatment with aerosol IFN-γ was well tolerated in a 2-year safety study in patients with idiopathic pulmonary fibrosis, where the decline in pulmonary function was reversed. The same approach may also be useful for COPD.82

Other types of drugs may also benefit from changing the mode of application. A drug that shows long term toxicity can still prove valuable when used for short periods of time. For example, mibefradil, originally used for blood pressure control, was withdrawn as chronic use slowed the excretion of as many as 26 other drugs, causing them to accumulate to dangerous levels in the liver. More recently, it was shown that short term dosing with mibefradil can enhance the activity of several different cancer drugs.52

2.3 | Targeting host cell pathways

Given the rising number of strains found to be resistant to drugs that affect pathways specific to infectious agents, many repurposing studies are now looking for more broad-spectrum agents that block or activate host pathways, such as those targeted in cancer therapy.13 Drugs developed to inhibit intracellular kinases,83,84 phosphatases85, or reduce inflammation may indeed have multivalent activity against several different infectious organisms. For example, drugs approved for different indications, chosen from a 400 compound library, inhibited the ability of several pathogenic bacteria to grow in macrophages. Trifluoperazine (antipsychotic), amoxapine (antidepressant), and doxapram (a breathing stimulant) protected up to 60% of animals tested against plague (Yersinia pestis), whereby full protection required coadministration of vancomycin.86 The same group found that, at 33 µM, 9% to 13% of 780 tested drugs inhibited the growth of Klebsiella pneumoniae or Acinetobacter baumannii in macrophages.

2.3.1 | Antiinflammatory agents

As noted above, the ability of thalidomide and its derivatives to block TNF-α, a major regulator of the inflammatory response, has led to their use in diseases such as psoriasis and other autoimmune disorders characterized by cytokine overproduction. The tetracycline derivative, doxycycline, can aid in wound healing through inhibiting host metalloproteases that may be causing further tissue injury.87,88 Doxycycline can also play a similar role in controlling sepsis.89

Antiinflammatory compounds, including nonsteroidal antiinflammatory drugs (NSAIDs), can control fever and other aspects of the “cytokine storm” that several different viruses induce. Ibuprofen has been suggested to control Ebola symptoms.90 NSAIDs can also improve survival in cancers, for example in patients with activating mutations in PIK3CA, which are very common in head and neck cancers. The predicted 5-year disease-specific and overall survival was significantly higher (72% vs 25%; 78% vs 45%, respectively) for those who used NSAIDs regularly than for those who did not.91
2.3.2 Stimulating antipathogen mechanisms

As treatment with agents targeting the pathogen directly can select for resistant viruses, repurposing of drugs designed to inhibit host cell pathways required by viruses or activate pathogen response mechanisms have been suggested as broad-spectrum antivirals. Recent epidemics have emphasized the need for wide spectrum oral agents to stimulate host antiviral pathways. While the earliest protein antivirals, interferons, can control many viruses, they are expensive, heat-sensitive, and even the new, extended half-life pegylated (IFN) preparations require regular injections. However, smaller antiviral compounds may have enhanced activity when combined with pegIFN. Small antivirals that induce the IFN pathway, identified through high throughput screening, such as Chugai’s RO8191 may provide a patient-friendly “oral interferon treatment” that could complement direct acting antivirals (DAAs). Alternatively, compounds that activate the STING pathway may also provide useful adjuvant treatments.

However, many viruses are not sensitive to the IFN response or produce factors that interfere with IFN and interferon-stimulated genes (ISGs). As IFN levels in uninfected individuals are typically vanishingly low, the high persistent presence of IFNs and/or their induced genes (ISGs) may also complicate or even cause other diseases, such as lupus or atopic dermatitis. Recent results indicate that high serum IFN- levels after treatment for HCV with DAAs may correlate with the development of hepatic carcinoma. Thus several IFN neutralizing therapies have been tested for lupus. These may be useful for treating other diseases characterized by overproduction of IFNs and their induced proteins. These include those that bind IFN itself, such as rontalizumab and inhibitors of interferon pathway proteins.

Autophagy pathways can be specifically targeted, according to the type of virus infection. Inhibitors of autophagy may inhibit the growth of RNA viruses that rely on the associated membranes for replication, such as Picornaviruses and Dengue. However, such inhibitors might favor the lytic growth of herpes and other viruses whose replication is held in check by autophagy.

Results of screening drug libraries have revealed several other mammalian cell pathways that may be targeted. For example, blockers of Ca2+ and other ion channels are reported to have in vitro activity against Ebola, Japanese Encephalitis virus, human cytomegalovirus and arenaviruses such as Lassa fever.

3 REPURPOSING AS A PATH TO NEW DRUGS

Simply having the desired activity does not mean that an older drug can move rapidly into the clinic. Older drugs may have significant problems that will interfere with their implementation in modern therapy programs, such as poor physicochemical properties, significant side effects or requiring unrealistic dosing for use in patients who are already severely ill. While repurposing an approved drug may seem to be the most rapid path to treatment, moving too rapidly into clinical testing can interfere with efforts to find more specific treatments. Extreme caution must be taken in identifying the best treatment for rare diseases, where the patient pool is small and fragile. Creutzfeldt-Jakob disease, as one example, is an extremely rare and rapidly moving syndrome. Many tests of repurposed compounds, including doxycycline, quinacrine, pentosan sulfate, and flupirtine, all failed to have any significant effect on patient survival, due to their lack of specificity.

Thus, to enhance specificity and deal with the issues discussed in Section 3.1, as well as licensing considerations, it may be best to regard that first active, approved drug as a starting point for developing a therapeutic. Collaboration with a medicinal chemistry group or company can promote access to derivatives with better specific activities, more suitable physicochemical properties, or reduced side effects.
3.1 | Obstacles to overcome in repurposing

3.1.1 | Many traditional and approved drugs have poor physicochemical properties

For example, fluorescein derivatives of rose bengal dye, approved for food use and a common laboratory stain, were first used medicinally to treat ocular infections in 1914. Thanks to its low toxicity, rose bengal therapies quickly entered the clinic. Today, topical versions of the dye are being tested for reducing scarring after injury or burns. The molecule became even more interesting when testing in the 1980s indicated it inhibited the growth of tumor cells. A formulation, PV-10, is in clinical trials for treating tumors. However, large amounts of the dye are required as the compound has only a 30-minute half-life in serum and as Figure 1 illustrates, it is a large hydrophobic molecule with limited solubility. The results of these trials may be used to obtain derivatives with better pharmaceutical characteristics, designed for solubility or more specific anticancer activities.

Other FDA-approved drugs are not ideal candidates for further development. Many chemotherapy drugs that require infusion, such as the bright blue compound mitoxantrone, an agent which has a variety of activities in cells, have limited solubility and may induce nonspecific aggregation of the protein or assay reagents in high throughput screening. A good deal of time can be lost to these nonspecific aggregators, which can have misleading activity in a variety of different assays at low micromolar concentrations. Mitoxantrone even turned up in a preliminary screen to identify inhibitors of uridylylation of the peptide linked to the genome, VPg, to VPgpU in the author’s group. Several different assays were required to show it precipitated the RNA substrate, rather than specifically inhibiting the polymerase. As with the uridylylation reaction, adding detergents to control aggregation induced by such agents may not be suitable for most assays. The physicochemical properties for drugs, such as solubility, can be obtained from free online databases, including ZINC or OCHEM (http://ochem.eu).

3.1.2 | Active compounds can have severe side effects

Many FDA-approved drugs, including those used for chemotherapy, have severe side effects that may prevent their use for other purposes. For example, emetine, better known as an active ingredient of ipecac syrup, binds ribosomes and inhibits a variety of disease-causing protozoa. It has also been reported to selectively kill acute myeloid leukemia cells. Emetine was selected by an Ebola minigenome HTS assay (which can be run at BSL-2 level) from a 2080 active component library and reported to have submicrometer activity against both Zika and Ebola virus.
However, emetine causes severe nausea when taken orally or injected. It remains to be seen whether emetine’s side effects can be overcome sufficiently to bring it into the clinic for these indications.

Similarly, the success of the many ongoing trials of calcineurin inhibitors (e.g., tacrolimus) for controlling inflammatory bowel disease, atopic dermatitis, rheumatoid arthritis, and autoimmune diseases may be limited by the increased risk of severe infection intrinsic to inhibiting a pathway that is essential for control of bacterial and fungal pathogens.146 Side effects may also limit the clinical usefulness of a variety of off-the (drug-)shelf compounds suggested as treatments for serious virus infections. The FDA-approved cancer drug, gemcitabine, inhibits picornaviruses147, and other viruses.148 The bioflavonoid rutin inhibits norovirus.149 Quinine and other antimalarials have been suggested for flaviviruses such as Dengue150,151 and Zika152 as well as the coronaviruses MERS/SARS.153 The clinical future of these treatments will depend on whether the side effects during short term administration are acceptable. It may be also possible to identify other known drugs with reduced side effects. A recent paper suggests that by categorizing drugs according to their indications and known side effects, one could identify drugs that could replace those with a “boxed warning” by safer ones with similar efficacy.154

3.1.3 | High effective concentrations may not be clinically achievable

Another obstacle to repurposing is that higher concentrations of the drug may be needed when used for purposes beyond the original design. High throughput screening for approved drugs may be conducted at concentrations of 33 to 100 µM, where compound screenings of larger compound libraries are done at 10 to 20 µM, followed by dilutions to lower concentrations in secondary assays of initial hits. At high concentrations, drugs may have different mechanisms than those associated with their suggested use in humans. Compounds may bind to off-target molecules that have little to do with the initially determined mechanism of actions, causing side effects when translated to human therapies. Even molecules suggested to be specific, for example, those with high affinity for a G-protein coupled receptor, have been found to bind completely unrelated molecules, such as phosphodiesterases, whereby the binding sites can be very different.155 Screening a large compound library by docking found many molecules selected to bind to a specific site in fact bound preferentially to a different one when given a larger region of the protein surface to choose from.136

The inexpensive, orally available nucleotide analogs, ribavirin, and favipiravir (developed for influenza156), when used at high concentrations, inhibit many viral RNA polymerases by different mechanisms.157-160 Favipiravir inhibits Congo hemorrhagic fever virus (CCHF)161 and Ebola162 in mice and Ebola163 and Lassa fever164 in primate models. It has been used with ribavirin to treat Lassa fever patients.165

However, favipiravir, even at doses of 150 to 300 mg/kg, did not completely eliminate CCHF in surviving infected mice, with several dying after treatment stopped. About 50% of the Ebola-infected macaques survived, at the highest dose used (180 mg/kg). Although favipiravir is a small and soluble compound (Figure 2), this is still a very high dose. When used in combination with oseltamivir (20 mg/kg) to treat influenza, effective doses were in the range of 50 mg/kg.166 Attempts to treat Ebola patients during the 2014 epidemic with high doses of favipiravir (1.2-2.4 g/day)162 gave unacceptable, significant side effects in patients who were more severely ill at presentation.167 Other nucleotide derivatives may have better potency in treating this disease,168 which is causing yet another large outbreak.

FIGURE 2  Favipiravir (T-705, avigan), a pyrazine carboxamide derivative, inhibits many different viral polymerases at high concentrations
In another example of a repurposed drug active only at high concentrations, the antimalarial drug pyrimethamine was found to stabilize β-hexosaminidase, the enzyme defective in Tay-Sachs disease. However, it produced significant side effects at the 75 to 100 mg doses required for efficacy in phases 1 and 2 testings in eight humans with late-onset Tay-Sachs. It was further tested more recently and gave a temporary improvement in measured enzyme activity, with one of four patients remaining stable while the other three continued to deteriorate. Seven years after the initial drug screening, more active forms of pyrimethamine were being sought, with the hope of finding a treatment for this deadly disease.

### TABLE 3 Tetracycline derivatives with different specificities

| Compound                | Structure                                                                 |
|-------------------------|---------------------------------------------------------------------------|
| Tetracycline            | ![](image1.png)                                                           |
| Doxycycline (metalo-protease inhibitor) | ![Doxycycline](image2.png)                                                |
| Omadacycline (nuzyra)   | ![Omadacycline](image3.png)                                               |
| Eravacycline (xerava)   | ![Eravacycline](image4.png)                                               |
| Sarecycline (seysara)   | ![Sarecycline](image5.png)                                                |

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### 3.2 Learning from thalidomide, designing better derivatives

Despite its worldwide withdrawal from the market in 1962, thalidomide was approved to treat granulomas associated with leprosy in the mid-1960s. This is an example of how, even with obvious side effects, a drug may be used clinically for a specific purpose. While thalidomide treatment will always be limited by its side effects, small changes resulted in much more active molecules that could be used at lower doses (Table 1). Of course, any derivative must go through toxicity screening and new clinical trials before it can be marketed, which can take considerable time. Thalidomide’s active derivatives were approved for specific treatments in 2005 (lenalidomide) and 2013 (pomalidomide).

Similarly, small structural changes may completely alter the spectrum of activity of an antibiotic. Conjugation of daptomycin, an antibiotic targeting Gram-positive bacteria, with an iron-binding siderophore mimetic generated a new series of antibiotics that could penetrate the cell wall of Gram-negative, antibiotic-resistant strains of Acinetobacter baumannii.

Recently approved derivatives of tetracycline (Table 3) also illustrate how small structural changes may enhance a drug’s specific activity or ability to evade resistance mechanisms. Tetracyclines block elongation of proteins in bacteria by binding to the 30S ribosomal subunit. New derivatives, such as Nuzyra (omadacycline), an aminomethylcycline, have been designed to overcome bacterial resistance to tetracycline, caused by bacterial efflux
pumps (tetK, tetL, tet) or ribosomal protection proteins such as tetM. Nuzyra can be used to treat bacterial pneumonia caused by a variety of resistant bacteria, including S. aureus, and streptococcal strains.\textsuperscript{174} Xerava (eravacycline), a synthetic fluorocycline, was developed to treat intra-abdominal infections caused by many different enteric bacteria.\textsuperscript{175,176} In contrast, Seysara (sarecycline), recommended for moderate-to-severe acne vulgaris, has less activity against enteric bacteria but reduces inflammation.\textsuperscript{177,178} Although treatment with these derivatives may save costs, compared with vancomycin,\textsuperscript{179} they are still much more expensive than their parent, tetracycline.

\section*{CONCLUSIONS}

Repurposing of older molecules can be a path to finding novel therapies. Screening approved drug libraries may be the most efficient way to identify treatments for rare and emerging diseases. Perhaps the best example of a repurposed drug is thalidomide, which despite its notorious side effects provides a valuable treatment for cancers and granulomas associated with leprosy. Table 2 illustrates how many other drugs are in testing for their usefulness in treating conditions beyond their original targets. Older antibiotics may also be repurposed by combining them with other active compounds, to achieve broad-spectrum activities against infectious agents.

However, many existing drugs have poor physicochemical properties and significant side effects when used at high concentrations. Identifying appropriate derivatives of an approved compound can, in the end, be the fastest path to the clinic. Derivatives of thalidomide and tetracycline provide excellent examples of how relatively small changes in a drug structure can greatly lower the dosage required and alter the spectrum of activity.

While by no means comprehensive, the many examples are shown here illustrate the great promise of the existing pharmacopeia and modifications thereof to treat “the many ills that flesh is heir to”.

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\section*{REFERENCES}

1. Marrugal-Lorenzo JA, Serna-Gallego A, González-González L, et al. Inhibition of adenovirus infection by mifepristone. \textit{Antiviral Res.} 2018;159:77-83.
2. Dyall J, Coleman CM, Hart BJ, et al. Repurposing of clinically developed drugs for treatment of Middle East respiratory syndrome coronavirus infection. \textit{Antimicrob Agents Chemother.} 2014;58(8):4885-4893.
3. Devillers J. Repurposing drugs for use against Zika virus infection. \textit{SAR QSAR Environ Res.} 2018;29(2):103-115.
4. Botting C, Kuhn RJ. Novel approaches to flavivirus drug discovery. \textit{Expert Opin Drug Discovery.} 2012;7(5):417-428.
5. Risse E, Nicoll AJ, Taylor WA, et al. Identification of a compound that disrupts binding of amyloid-β to the prion protein using a novel fluorescence-based assay. \textit{J Biol Chem.} 2015;290(27):17020-17028.
6. Mosheri H, Mehta V, Yip CW, Salavati R. Pilot-scale compound screening against RNA editing identifies trypanocidal agents. \textit{J Biomol Screening.} 2015;20(1):92-100.
7. Rzuczek SG, Southern MR, Disney MD. Studying a drug-like, RNA-focused small molecule library identifies compounds that inhibit RNA toxicity in myotonic dystrophy. \textit{ACS Chem Biol.} 2015;10(12):2706-2715.
8. Corsello SM, Bittker JA, Liu Z, et al. The drug repurposing hub: a next-generation drug library and information resource. \textit{Nature Med.} 2017;23(4):405-408.
9. Shameer K, Glicksberg BS, Hodos R, et al. Systematic analyses of drugs and disease indications in RepurposeDB reveal pharmacological, biological and epidemiological factors influencing drug repositioning. Briefings in Bioinform. 2018;19(4):656-678.

10. Moosavinasab S, Patterson J, Strouse R, et al. RE:fine drugs: an interactive dashboard to access drug repurposing opportunities. Database. 2016;2016:baw083.

11. Brown AS, Patel CJ. A standard database for drug repositioning. Sci Data. 2017;4:170029.

12. Bao J, Marathe B, Govorkova EA, Zheng JJ. Drug repurposing identifies inhibitors of oseltamivir-resistant influenza viruses. Angew Chem, Int Ed. 2016;55(10):3438-3441.

13. Gupta SC, Sung B, Prasad S, Webb LJ, Aggarwal BB. Cancer drug discovery by repurposing: teaching new tricks to old dogs. Trends Pharmacol Sci. 2013;34(9):508-517.

14. Bonine NG, Berger A, Altmancat A, et al. Impact of delayed appropriate antibiotic therapy on patient outcomes by antibiotic resistance status from serious gram-negative bacterial infections. Am J Med Sci. 2019;357(2):103-110.

15. Chen D, Misra M, Sower L, Peterson JW, Kellogg GE, Schein CH. Novel inhibitors of anthrax edema factor. Bioorg Med Chem. 2008;16(15):7225-7233.

16. Schein CH, Chen D, Ma L, et al. Pharmacophore selection and redesign of nonnucleotide inhibitors of anthrax edema factor. Toxins. 2012;4(11):1288-1300.

17. Johansen LM, DeWald LE, Shoemaker CJ, et al. A screen of approved drugs and molecular probes identifies therapeutics with anti-Ebola virus activity. Sci Transl Med. 2015;7(290):290ra89-290ra89.

18. Perwitasari O, Yan X, O'Donnell J, Johnson S, Tripp RA. Repurposing kinase inhibitors as antiviral agents to control influenza A virus replication. Assay Drug Dev Technol. 2015;13(10):638-649.

19. Rayasam GV, Balganesh TS. Exploring the potential of adjunct therapy in tuberculosis. Trends Pharmacol Sci. 2015;36(8):506-513.

20. Kozbial K, Moser A, Al-Zaairy R, et al. Follow-up of sustained virological responders with hepatitis C and advanced liver disease after interferon/ribavirin-free treatment. Liver Int. 2018;38(6):1028-1035.

21. Meanwell NA. Improving drug design: an update on recent applications of efficiency metrics, strategies for replacing problematic elements, and compounds in nontraditional drug space. Chem Res Toxicol. 2016;29(4):564-616.

22. Rehman W, Arfons LM, Lazarus HM. The rise, fall and subsequent triumph of thalidomide: lessons learned in drug development. Ther Adv Hematol. 2011;2(5):291-308.

23. Kelsey FO. Thalidomide update: regulatory aspects. Teratology. 1988;38(3):221-226.

24. Mori T, Ito T, Liu S, et al. Structural basis of thalidomide enantiomer binding to cereblon. Sci Rep. 2018;8(1):1294.

25. Zhu YX, Kortuem KM, Stewart AK. Molecular mechanism of action of immune-modulatory drugs thalidomide, lenalidomide and pomalidomide in multiple myeloma. Leukemia & lymphoma. 2013;54(4):683-687.

26. Martinucci F, Giovinazzo J, Tan AU, et al. Lessons of leprosy: the emergence of TH17 cytokines during type II reactions (ENL) is teaching us about T-cell plasticity. J Drugs Dermatol. 2012;11(5):626-630.

27. Petzold G, Fischer ES, Thomä NH. Structural basis of lenalidomide-induced CK1α degradation by the CRL4CRBN ubiquitin ligase. Nature. 2016;532(7597):127-130.

28. Scribano ML, Cantoro L, Marullo M, Cosintino R, Kohn A. Mucosal healing with thalidomide in refractory Crohn's disease patients intolerant of anti-TNF-α drugs: report of 3 cases and literature review. J Clin Gastroenterol. 2014;48(6):530-533.

29. Lagier JC, Raoult D. Immune reconstitution inflammatory syndrome associated with bacterial infections. Expert Opin Drug Saf. 2014;13(3):341-350.

30. Starke RM, Chalouhi N, Jabbour PM, et al. Critical role of TNF-α in cerebral aneurysm formation and progression to rupture. J Neuroinflammation. 2014;11:77.

31. Batsaikhans B, Wang JY, Scerba M, et al. Postinjury neuroprotective effects of the thalidomide analog 3,6'-dithiothalidomide on traumatic brain injury. Int J Mol Sci. 2019;20(3):502.

32. Corral LG, Kaplan G. Immunomodulation by thalidomide and thalidomide analogues. Ann Rheum Dis. 1999;58(Suppl 1):i107-i113.

33. Corral LG, Haslett PA, Muller GW, et al. Differential cytokine modulation and T cell activation by two distinct classes of thalidomide analogues that are potent inhibitors of TNF-alpha. J Immunol (Baltimore, Md.: 1950). 1999;163(1):380-386.

34. De Souza A, Strober BE, Merola JF, Oliver S, Franks AG Jr Jr. Apremilast for discoid lupus erythematosus: results of a phase 2, open-label, single-arm, pilot study. J Drugs Dermatol. 2012;11(10):1224-1226.

35. Dasanu CA, Mitzis D, Alexandrescu DT. Hair repigmentation associated with the use of lenalidomide: graying may not be an irreversible process! J Oncol Pharm Pract. 2013;19(2):165-169.

36. Ono S, Miyachi Y, Arakawa A. Hair regrowth following TNF-α blockade in coexisting psoriasis vulgaris and alopecia areata. European journal of dermatology: EJD. 2013;23(4):537.
37. Laurikkala J, Pispä J, Jung HS, et al. Regulation of hair follicle development by the TNF signal ectodysplasin and its receptor Edar. Development. 2002;129(10):2541-2553.
38. Robert C, Spatz A, Faivre S, Armand JP, Raymond E. Tyrosine kinase inhibition and grey hair. The Lancet. 2003;361(9362):1056.
39. Rivera N, Boada A, Bielsa MI, et al. Hair repigmentation during immunotherapy treatment with an antiprogrammed cell death 1 and antiprogrammed cell death ligand 1 agent for lung cancer. JAMA Dermatol. 2017;153(11):1162-1165.
40. Polster SP, Stadnik A, Akers AL, et al. Atorvastatin treatment of cavernous angiomas with symptomatic hemorrhage: exploratory proof of concept (AT CASH EPOC) trial. Neurosurgery. 2018. (in press).
41. Abdullah MI, de Wolf E, Jawad MJ, Richardson A. The poor design of clinical trials of statins in oncology may explain their failure—lessons for drug repurposing. Cancer Treat Rev. 2018;69:84-89.
42. Walker VM, Davies NM, Jones T, Kehoe PG, Martin RM. Can commonly prescribed drugs be repurposed for the prevention or treatment of Alzheimer’s and other neurodegenerative diseases? Protocol for an observational cohort study in the UK Clinical Practice Research Datalink. BMJ Open. 2016;6(12):e012044.
43. Rowbotham SE, Krishna SM, Moran CS, Golledge J. Fenofibrate and telmisartan in the management of abdominal aortic aneurysm. Curr Drug Targets. 2018;19(11):1241-1246.
44. Connick P, De Angelis F, Parker RA, et al. Multiple sclerosis-secondary progressive multiarm randomisation trial (MS-SMART): a multiarm phase Ibb randomised, double-blind, placebo-controlled clinical trial comparing the efficacy of three neuroprotective drugs in secondary progressive multiple sclerosis. BMJ Open. 2018;8(8):e021944.
45. Eleuteri C, Olla S, Veroni C, et al. A staged screening of registered drugs highlights remyelinating drug candidates for clinical trials. Sci Rep. 2017;7:45780.
46. Panch SR, Bozik ME, Brown T, et al. Dexamethasone as an oral steroid-sparing agent in hypereosinophilic syndromes. Blood. 2018;132(5):501-509.
47. Reinhart S, Stoye N, Luderer M, et al. Identification of disulfiram as a secretase-modulating compound with beneficial effects on Alzheimer’s disease hallmarks. Sci Rep. 2018;8(1):1329.
48. Parikh AB, Kozuch P, Rohs N, Becker DJ, Levy BP. Metformin as a repurposed therapy in advanced non-small cell lung cancer (NSCLC): results of a phase II trial. Invest New Drugs. 2017;35(6):813-819.
49. Long DE, Peck BD, Martz JL, et al. Metformin to augment strength training effective response in seniors (MASTERS): study protocol for a randomized controlled trial. Trials. 2017;18(1):192.
50. Ran Y, Hossain F, Pannuti A, et al. γ-Secretase inhibitors in cancer clinical trials are pharmacologically and functionally distinct. EMBO Mol Med. 2017;9(7):950-966.
51. Nygaard HB, Wagner AF, Bowen GS, et al. A phase Ib multiple ascending dose study of the safety, tolerability, and central nervous system availability of AZD0530 (saracatinib) in Alzheimer’s disease. Alzheimer’s Research & Therapy. 2015;7(1):35.
52. Krouse AJ, Gray L, Macdonald T, McCray J. Repurposing and Rescuing of Migbefradil, an antihypertensive, for cancer: a case study. Assay Drug Dev Technol. 2015;13(10):650-653.
53. Blumenthal GM, Gills JJ, Ballas MS, et al. A phase I trial of the HIV protease inhibitor nelfinavir in adults with solid tumors. Oncotarget. 2014;5(18):18116-18172.
54. China L, Skene SS, Bennett K, et al. ATTIRE: albumin To prevenT infection in chronic liveR failurE: study protocol for an interventional randomised controlled trial. BMJ Open. 2018;8(10):e023754.
55. Masaki C, Sharpley AL, Godlewksa BR, et al. Effects of the potential lithium-mimetic, ebsealen, on brain neurochemistry: a magnetic resonance spectroscopy study at 7 tesla. Psychopharmacology. 2016;233(6):1097-1104.
56. Hellings JA, Reed G, Cain SE, et al. Loxapine add-on for adolescents and adults with autism spectrum disorders and irritability. J Child Adolesc Psychopharmacol. 2015;25(2):150-159.
57. Rossignol JF. Nitazoxanide: a first-in-class broad-spectrum antiviral agent. Antiviral Res. 2014;110:94-103.
58. Bakker AC, La Rosa S, Sherman LS, et al. Neurofibromatosis as a gateway to better treatment for a variety of malignancies. Prog Neurobiol. 2017;152:149-165.
59. Bumb JM, Enning F, Leweke FM. Drug repurposing and emerging adjunctive treatments for schizophrenia. Expert Opin Pharmacother. 2015;16(7):1049-1067.
60. Tyers M, Wright GD. Drug combinations: a strategy to extend the life of antibiotics in the 21st century. Nat Rev Microbiol. 2019;17:141-155.
61. Hartkoorn RC, Sala C, Neres J, et al. Towards a new tuberculosis drug: pyrazinamide—nature’s isoniazid. EMBO Mol Med. 2012;4(10):1032-1042.
62. Diacon AH, Dawson R, von Groote-Bidlingmaier F, et al. Bactericidal activity of pyrazinamide and clofazimine alone and in combinations with pretomanid and bedaquiline. Am J Respir Crit Care Med. 2015;191(8):943-953.
63. Rotondo CM, Wright GD. Inhibitors of metallo-β-lactamases. Curr Opin Microbiol. 2017;39:96-105.
64. Wright GD. Antibiotic adjuvants: rescuing antibiotics from resistance. Trends Microbiol. 2016;24(11):862-871.
93. Weissmann C, Nagata S, Boll W, et al. Structure and Expression of Human Alpha-interferon Genes. *Interferons*. 25. Academic Press; 1982:295-326.
94. Weissmann C, Nagata S, Boll W, et al. Structure and expression of human IFN-alpha genes. *Philos Trans R Soc B*. 1982;299(1094):7-28.
95. Schoggins JW. Recent advances in antiviral interferon-stimulated gene biology. *F1000Research*. 2018;7:309.
96. Maughan A, Ogbaru O. Pegylated interferon alpha 2a for the treatment of hepatitis C virus infection. *Expert Opin Drug Metab Toxicol*. 2018;14(2):219-227.
97. Hagiwara S, Nishida N, Watanabe T, et al. Sustained antiviral effects and clearance of hepatitis surface antigen after combination therapy with entecavir and pegylated interferon in chronic hepatitis B. *Antivir Ther*. 2018;23(6):513-521.
98. Konishi H, Okamoto K, Ohmori Y, et al. An orally available, small-molecule interferon inhibits viral replication. *Sci Rep*. 2012;2:259.
99. Cerón S, North BJ, Taylor SA, Leib DA. The STING agonist 5,6-dimethylxanthenone-4-acetic acid (DMXAA) stimulates an antiviral state and protects mice against herpes simplex virus-induced neurological disease. *Virology*. 2019;529:23-28.
100. Skouboe MK, Knudsen A, Reinert LS, et al. STING agonists enable antiviral cross-talk between human cells and confer protection against genital herpes in mice. *PloS Pathog*. 2018;14(4). e1006976.
101. Mutz P, Metz P, Lempp FA, et al. HBV bypasses the innate immune response and does not protect HCV from antiviral activity of interferon. *Gastroenterology*. 2018;154(6):1791-1804. e1722.
102. Liu R, Moss B. Vaccinia virus C9 ankyrin repeat/F-box protein is a newly identified antagonist of the type I interferon-induced antiviral state. *J Virol*. 2018;92(9):e00053-00118. https://doi.org/1128/JVI.00053-18
103. Kalinowski A, Galen BT, Ueki IF, et al. Respiratory syncytial virus activates epidermal growth factor receptor to suppress interferon regulatory factor 1-dependent interferon-lambda and antiviral defense in airway epithelium. *Mucosal Immunol*. 2018;11(3):958-967.
104. Feng L, Sheng J, Yu GP, et al. Human cytomegalovirus UL23 inhibits transcription of interferon-γ stimulated genes and blocks antiviral interferon-γ responses by interacting with human N-myc interactor protein. *PloS Pathog*. 2018;14(1):e1006867.
105. Mok CC. Biological and targeted therapies of systemic lupus erythematosus: evidence and the state of the art. *Expert Rev Clin Immunol*. 2017;13:677-692.
106. Inoue-Shinomiya E, Murakawa M, Asahina Y, et al. Association of serum interferon-λ3 levels with hepatocarcinogenesis in chronic hepatitis C patients treated with direct-acting antiviral agents. *Hepatol Res*. 2019;49:500-511.
107. Baker KF, Isaacs JD. Novel therapies for immune-mediated inflammatory diseases: what can we learn from their use in rheumatoid arthritis, spondyloarthritis, systemic lupus erythematosus, psoriasis, Crohn’s disease and ulcerative colitis? *Ann Rheum Dis*. 2018;77(2):175-187.
108. Maurer B, Bosanac I, Shia S, et al. Structural basis of the broadly neutralizing anti-interferon-α antibody rontalizumab: neutralizing anti-interferon-α antibody rontalizumab. *Protein Sci*. 2015;24(9):1440-1450.
109. Khamashta M, Merrill JT, Werth VP, et al. Sifalimumab, an anti-interferon-α monoclonal antibody, in moderate to severe systemic lupus erythematosus: a randomised, double-blind, placebo-controlled study. *Ann Rheum Dis*. 2016;75(11):1909-1916.
110. Miller S, Tsou PS, Coit P, et al. Hypomethylation of STAT1 and HLA-DRB1 expression in lupus CD8+ T cells. *Ann Rheum Dis*. 2019;78:519-528.
111. Zimmermann N, Wolf C, Schwenke R, et al. Assessment of clinical response to janus kinase inhibition in patients with familial chilblain lupus and TREX1 mutation. *JAMA dermatology*. 2019;155:342.
112. Kahn JS, Deverapalli SC, Rosmarin DM. JAK-STAT signaling pathway inhibition: a role for treatment of discoid lupus erythematosus and dermatomyositis. *Int J Dermatol*. 2018;57(8):1007-1014.
113. Brown HM, Biering SB, Zhu A, Choi J, Hwang S. Demarcation of viral shelters results in destruction by membrane-bound GTPases: antiviral function of autophagy proteins and interferon-inducible GTPases. *BioEssays*. 2018;40(6):1700231.
114. Subramanian G, Kuzmanovic T, Zhang Y, et al. A new mechanism of interferon’s antiviral action: Induction of autophagy, essential for paramyxovirus replication, is inhibited by the interferon stimulated gene, TDRD7. *PloS Pathog*. 2018;14(1):e1006877.
115. Liu Y, Cherry S. Zika virus infection activates sting-dependent antiviral autophagy in the Drosophila brain. *Autophagy*. 2019;15(1):174-175.
116. Kim SR, Song JH, Ahn JH, et al. Antiviral and anti-inflammatory activity of budesonide against human rhinovirus infection mediated via autophagy activation. *Antiviral Res*. 2018;151:87-96.
117. Delorme-Axford E, Abernathy E, Lennemann NJ, et al. The exoribonuclease Xrn1 is a post-transcriptional negative regulator of autophagy. *Autophagy*. 2018;14(5):898-912.
118. Mateo R, Nagamine CM, Spagnolo J, et al. Inhibition of cellular autophagy deranges dengue virion maturation. *J Virol*. 2013;87(3):1312-1321.
119. Schein CH. Polyglutamine repeats in viruses. *Mol Neurobiol*. 2019;56:3664-3675.
120. Yin Y, Deng W, Zhou X, et al. PI3K-Akt-mTOR axis sustains rotavirus infection via the 4E-BP1 mediated autophagy pathway and represents an antiviral target. *Virulence*. 2018;9(1):83-98.

121. Dyall J, Johnson JC, Hart BJ, et al. In vitro and in vivo activity of amiodarone against Ebola virus. *J Infect Dis*. 2018;218:5592-5596.

122. Wang S, Liu Y, Guo J, et al. Screening of FDA-approved drugs for inhibitors of Japanese Encephalitis virus infection. *J Virol*. 2017;91(21):e01055-e01117. https://doi.org/10.1128/JVI.01055-17

123. Mercorelli B, Luganini A, Celegato M, Palù G, Gribaudo G, Lorean GiA. Repurposing the clinically approved calcium antagonist manidipine dihydrochloride as a new early inhibitor of human cytomegalovirus targeting the Immediate-Early 2 (IE2) protein. *Antiviral Res*. 2018;150:130-136.

124. Wang P, Liu Y, Zhang G, et al. Screening and Identification of Lassa virus entry inhibitors from an FDA-approved drug library. *J Virol*. 2018;92(16):e00954-e01018. https://doi.org/10.1128/JVI.00954-18 [published online ahead of print August 15, 2018].

125. Buranda T, Gineste C, Wu Y, et al. A high-throughput flow cytometry screen identifies molecules that inhibit hantavirus cell entry. *SLAS discov. 2018;247:255521876623.

126. Mead S, Tagliavini F. Handbook of clinical neurology. *Clinical Trials*. 2018;153:431-444.

127. Hernandez JJ, Przyzak M, Smith L, et al. Giving drugs a second chance: overcoming regulatory and financial hurdles in repurposing approved drugs as cancer therapeutics. *Front Oncol*. 2017;7:273.

128. Patil N, Truong V, Holmberg MH, et al. Safety and efficacy of rose bengal derivatives for glial scar ablation in chronic spinal cord injury. *J Neurotrauma*. 2018;35(15):1745-1754.

129. Read TA, Smith A, Thomas J, et al. Intralesion PV-10 for the treatment of in-transit melanoma metastases-results of a prospective, nonrandomized, single center study. *J Surg Oncol*. 2018;117(4):579-587.

130. Dahlin JL, Nissink JWM, Francis S, et al. Post-HTS case report and structural alert: promiscuous 4-aroyl-1,5-disubstituted-3-hydroxy-2H-pyrrl-2-one actives verified by ALARM NMR. *Bioorg Med Chem Lett*. 2015;25(21):4740-4752.

131. Wu KJ, Zhong HJ, Li G, et al. Structure-based identification of a NEDD8-activating enzyme inhibitor via drug repurposing. *Eur J Med Chem*. 2018;143:1021-1027.

132. Baell JB, Nissink JWM. Seven year itch: Pan-Assay Interference Compounds (PAINS) in 2017—utility and limitations. *ACS Chem Biol*. 2018;13(1):36-44.

133. Schein CH, Volk DE, Oezguen N, Novel P. Structure-based mechanism for uridylylation of the genome-linked peptide (VPg) of picornaviruses. *Proteins*. 2006;63(4):719-726.

134. Schein CH, Oezguen N, van der Heden van Noort GJ, et al. NMR solution structure of poliovirus uridylylated peptide linked to the genome (VPgpU). *Peptides*. 2010;31(8):1441-1448.

135. Schein CH, Ye M, Paul AV, et al. Sequence specificity for uridylylation of the viral peptide linked to the genome (VPg) of enteroviruses. *Virology*. 2015;484:80-85.

136. Schein CH, Rowold D, Choi KH. Allosteric inhibitors of Coxsackie virus A24 RNA polymerase. *Bioorg Med Chem*. 2016;24(4):570-577.

137. Sterling T, Irwin JJ. ZINC 15—ligand discovery for everyone. *J Chem Inf Model*. 2015;55(11):2324-2337.

138. Sushko I, Novotarskyy S, Körner R, et al. Online chemical modeling environment (OCHEM): web platform for data storage, model development and publishing of chemical information. *J Comput Aided Mol Des*. 2011;25(6):533-554.

139. Wong W, Bai X, Brown A, et al. Cryo-EM structure of the Plasmodium falciparum 80S ribosome bound to the anti-protozoan drug emetine. *eLife*. 2014;3:3. https://doi.org/10.7554/eLife.03080

140. Matthews H, Deakin J, Rajab M, Idris-Usem M, Nirmalan NJ. Investigating antimalarial drug interactions of emetine dihydrochloride hydrate using CalcuSyn-based interactivity calculations. *PLoS One*. 2017;12(3):e0173303.

141. Krstin S, Mohamed T, Wang X, Wink M. How do the alkaloids emetine and homoharringtonine kill trypanosomes? An insight into their molecular modes of action. *Phytochemistry*. 2016;23(14):1771-1777.

142. Mirza H, Teo JDW, Upcroft J, Tan KSW. A rapid, high-throughput viability assay for Blastocystis spp. reveals metronidazole resistance and extensive subtype-dependent variations in drug susceptibilities. *Antimicrob Agents Chemother*. 2011;55(2):637-648.

143. Prabhu R, Sehgal R, Chakraborti A, Malla N, Ganguly UK, Mahajan RC. Isolation of emetine resistant clones of *Entamoeba histolytica* by petri dish agar method. *Indian J Med Res*. 2000;111:11-13.

144. Cornet-Masana JM, Moreno-Martinez D, Lara-Castillo MC, et al. Emetine induces chemosensitivity and reduces clonogenicity of acute myeloid leukemia cells. *Oncotarget*. 2016;7(17):23239-23250.

145. Edwards MR, Pietzsch C, Vausselin T, Shaw ML, Bukreyev A, Basler CF. High-Throughput minigenome system for identifying small-molecule inhibitors of ebola virus replication. *ACS Infect Dis*. 2015;1(8):380-387.

146. Bendickova K, Tidu F, Fric J. Calcineurin-NFAT signalling in myeloid leucocytes: new prospects and pitfalls in immunosuppressive therapy. *EMBO Mol Med*. 2017;9(8):990-999.
147. Zhang Z, Yang E, Hu C, et al. Cell-based high-throughput screening assay identifies 2′,2′-difluoro-2′-deoxycytidine gemcitabine as a potential antipoliovirus agent. *ACS Infect. Dis.* 2017;3(1):45-53.

148. Lee K, Kim DE, Jang KS, Kim SJ, Cho S, Kim C. Gemcitabine, a broad-spectrum antiviral drug, suppresses enterovirus infections through innate immunity induced by the inhibition of pyrimidine biosynthesis and nucleotide depletion. *Onctarget.* 2017;8(70):115315-115235.

149. Chéron N, Yu C, Kolawaole AO, Shakhnovich EI, Wobus CE. Repurposing of rutin for the inhibition of norovirus replication. *Arch Virol.* 2015;160(9):2353-2358.

150. Malakar S, Sreelatha L, Dechtawewat T, et al. Drug repurposing of quinine as antiviral against dengue virus infection. *Drug Repurposing of Quinine as Antiviral Against Dengue Virus Infection.* 2018;255:171-178.

151. Han Y, Mesplède T, Xu H, Quan Y, Wainberg MA. The antimalarial drug amodiaquine possesses anti-ZIKA virus activities. *J Med Virol.* 2018;90(5):796-802.

152. Boonyasuppayakorn S, Reichert ED, Manzano M, Nagarajan K, Padmanabhan R. Amodiaquine, an antimalarial drug, inhibits dengue virus type 2 replication and infectivity. *Antiviral Res.* 2014;106:125-134.

153. Furuta Y, Komeno T, Nakamura T Favipiravir (T-705), a broad spectrum inhibitor of viral RNA polymerase. *Antiviral Res.* 2013:100(2):679-691.

154. Gowen BB, Westover JB, Sefing EJ, et al. Enhanced protection against experimental Junin virus infection through the use of a modified favipiravir loading dose strategy. *Antiviral Res.* 2017:145:131-135.

155. Baretier S, Sterling T, O’Meara MJ, Shoichet BK. The recognition of identical ligands by unrelated proteins. *ACS Chem. Biol.* 2015;10(12):2772-2784.

156. Naqvi S, Gohar MS, Qamar M, et al. Antiviral efficacy of favipiravir against Ebola virus: a translational study in cynomolgus macaques. *Antiviral Res.* 2017;145:131-135.

157. Baz M, Carbonneau J, Rhéaume C, Cavanagh MH, Boivin G. Combination therapy with oseltamivir and favipiravir delays mortality but does not prevent oseltamivir resistance in immunodeficient mice infected with pandemic A (H1N1) Influenza virus. *Viruses.* 2018;10(11):454.

158. Ngoc TT, Guo Q, McMillen B, et al. Investigating drug repositioning opportunities in FDA drug labels through topic modeling. *BMC Bioinformatics.* 2012:13(Suppl 15):S6.

159. Chaliha GC, D’Souza M, Gudimetla S, et al. Middle east respiratory syndrome and severe acute respiratory syndrome: current therapeutic options and potential targets for novel therapies. *Drugs.* 2017;77(18):1935-1966.

160. Bisgin H, Liu Z, Kelly R, Fang H, Xu X, Tong W. Investigating drug repositioning opportunities in FDA drug labels by similarity searching. *Oncotarget.* 2018;9(70):115315-115325.

161. Dyall J, Gross R, Kindrachuk J, et al. Middle east respiratory syndrome and severe acute respiratory syndrome: current therapeutic options and potential targets for novel therapies. *Drugs.* 2017;77(18):1935-1966.

162. Han Y, Mesplède T, Xu H, Quan Y, Wainberg MA. The antimalarial drug amodiaquine possesses anti-ZIKA virus activities. *J Med Virol.* 2018;90(5):796-802.

163. Furuta Y, Komeno T, Nakamura T Favipiravir (T-705), a broad spectrum inhibitor of viral RNA polymerase. *Antiviral Res.* 2013:100(2):679-691.

164. Vanharen J, Hout Vrancken B, Van Houdt J, et al. Distinct effects of T-705 (Favipiravir) and Ribavirin on Influenza virus replication and viral RNA synthesis. *Antimicrob Agents Chemother.* 2016;60(11):6679-6691.

165. Furuta Y, Takahashi K, Shiraki K, Smee DF, Barnard DL. Favipiravir (T-705), a novel viral RNA polymerase inhibitor. *Antiviral Res.* 2013;100(2):679-691.

166. Gowen BB, Westover JB, Sefing EJ, et al. Enhanced protection against experimental Junin virus infection through the use of a modified favipiravir loading dose strategy. *Antiviral Res.* 2017:145:131-135.

167. Han Y, Mesplède T, Xu H, Quan Y, Wainberg MA. The antimalarial drug amodiaquine possesses anti-ZIKA virus activities. *J Med Virol.* 2018;90(5):796-802.

168. Dyall J, Gross R, Kindrachuk J, et al. Middle east respiratory syndrome and severe acute respiratory syndrome: current therapeutic options and potential targets for novel therapies. *Drugs.* 2017;77(18):1935-1966.

169. Bisgin H, Liu Z, Kelly R, Fang H, Xu X, Tong W. Investigating drug repositioning opportunities in FDA drug labels through topic modeling. *BMC Bioinformatics.* 2012:13(Suppl 15):S6.

170. Barelier S, Sterling T, O’Meara MJ, Shoichet BK. The recognition of identical ligands by unrelated proteins. *ACS Chem. Biol.* 2015;10(12):2772-2784.

171. Osher E, Fattal FEBS J. 2012;2784. 2018;255:171-178.

172. Tropak MB, Mahuran D. Lending a helping hand, screening chemical libraries for compounds that enhance β-hexosaminidase activity in GM2 gangliosidosis cells: screening libraries for hexosaminidase enhancers. *FEBS J.* 2007:274(19):4951-4961.

173. Clarke JTR, Mahuran DJ, Sathe S, et al. An open-label Phase I/II clinical trial of pyrimethamine for the treatment of patients affected with chronic GM2 gangliosidosis (Tay-Sachs or Sandhoff variants). *Mol Gen Metab.* 2011;102(1):6-12.

174. Osher E, Fattal-Valevski A, Sagie L, et al. Effect of cyclic, low dose pyrimethamine treatment in patients with late onset Tay Sachs: an open label, extended pilot study. *Orphanet J Rare Dis.* 2015;10:45.

175. Tropak MB, Zhang J, Yonekawa S, et al. Pyrimethamine derivatives: insight into binding mechanism and improved enhancement of mutant β-N-acetylhexosaminidase activity. *J Med Chem.* 2015;58(11):4483-4493.

176. Ghosh M, Lin YM, Miller PA, Möllmann U, Boggess WC, Miller MJ. Siderophore conjugates of daptomycin are potent inhibitors of carbapenem resistant strains of acinetobacter baumannii. *ACS Infect. Dis.* 2018;4(10):1529-1535.
174. Barber KE, Bell AM, Wingler MJB, Wagner JL, Stover KR. Omadacycline enters the ring: a new antimicrobial contender. *Pharmacotherapy*. 2018;38(12):1194-1204.
175. Zhanel GG, Cheung D, Adam H, et al. Review of eravacycline, a novel fluorocycline antibacterial agent. *Drugs*. 2016;76(5):567-588.
176. Seifert H, Stefanik D, Sutcliffe JA, Higgins PG. In-vitro activity of the novel fluorocycline eravacycline against carbapenem nonsusceptible *Acinetobacter baumannii*. *Int J Antimicro Ag*. 2018;51(1):62-64.
177. Zhanel G, Critchley I, Lin LY, Alvandi N. Microbiological profile of sarecycline, a novel targeted spectrum tetracycline for the treatment of acne vulgaris. *Antimicrob Agents Chemother*. 2018;63(1):e01297-e01318. https://doi.org/10.1128/AAC.01297-18 [published online ahead of print January, 2019].
178. Moore A, Green LJ, Bruce S, et al. Once-daily oral sarecycline 1.5 mg/kg/day is effective for moderate to severe acne vulgaris: results from two identically designed, phase 3, randomized, double-blind clinical trials. *J Drugs Dermatol: JDD*. 2018;17(9):987-996.
179. LaPensee K, Lodise T. Potential cost-savings with once-daily aminomethylcycline antibiotic versus vancomycin in hospitalized patients with acute bacterial skin and skin structure infections. *Am Health Drug Benefits*. 2018;11(9):449-459.

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