Emergence of Third Generation Tetracyclines: New Magic Bullets to Tackle Antibiotic Resistance in the Post-antibiotic Era

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Authors’ contributions

This work was carried out in collaboration among all authors. All authors contributed to the review design and plan. Authors PJ and SG contributed to the data search, collection, extraction, and quality assessment for this review. Author SSVP created the figure and author PJ created the tables for the manuscript. All authors wrote the text, reviewed and edited the manuscript, and made substantial contributions to discussions of the content. All authors read and approved the final manuscript.

ABSTRACT

Tetracyclines are broad-spectrum antibiotics effective against a wide variety of microorganisms including bacteria, both Gram-positive and Gram-negative, mycoplasmas, rickettsiae, chlamydiae, and protozoan parasites. Owing to their broad-spectrum antimicrobial activities and inexpensiveness, tetracyclines have been used extensively in both human and animal infections and in animal feed as growth promoters. Owing to this, the global prevalence of antibiotic
resistance, particularly of tetracyclines, for Gram-positive methicillin-resistant Staphylococcus aureus (MRSA) and Streptococcus pneumoniae and Gram-negative extended-spectrum β-lactamase-producing Escherichia coli and Klebsiella species is high. Indeed, the acquisition of tetracycline-specific resistance genes, mutations within the ribosomal binding site, and/or chromosomal mutations lead to both class-specific and intrinsic antimicrobial resistance (AMR) mechanisms. As drug resistance increases globally, rendering diseases difficult to manage and eventually to mortality, antibiotics are becoming progressively less effective. Therefore, the discovery and development of novel antibiotics with appropriate indications is of the utmost importance. Among all antibiotics, the tetracyclines have acquired much attention due to the optimization of their chemical structures that paved the way to develop and introduce modern tetracyclines, referred to as third generation, namely tigecycline in the recent past, followed by omadacycline, eravacycline, and sarecycline very recently. Intriguingly, these novel tetracyclines are unique in two ways, first, these are highly effective against pathogens that acquired tetracycline-class resistance, and second, these agents exhibit either narrow or broad spectrum of in vitro activity against Gram-positive, Gram-negative, anaerobic, and atypical pathogens, including many drug-resistant strains lead to approval for limited use and unique indications. These beneficial effects represent a new era in the rational use of newer tetracyclines, the new magic bullets to tackle AMR in the post-antibiotic era. The present review focuses on third-generation tetracyclines emphasizing their safety, efficacy, and therapeutic choices in various clinical conditions of in-patient and out-patient settings.

Keywords: Antimicrobial resistance; eravacycline; multidrug resistance; omadacycline; sarecycline; tigecycline; tetracyclines; third generation.

1. INTRODUCTION

Pre-antibiotic era is the period before the discovery and potential use of the first antibiotic wherein the understanding and knowledge about microbes and infectious diseases were inadequate. During this period, epidemics, morbidity, and mortality due to infectious diseases were common due to the lack of successful approaches for treatment and prevention of the spread of contagious diseases [1,2]. Paul Ehrlich’s discovery of salvarsan in 1909 for the treatment of syphilis led to the basis of the concept of chemotherapy. This chemical, also known as compound 606 or arsphenamine, is widely termed the first magic bullet [2-4]. “Several chemicals with disinfectant and bacteriostatic potential have been developed and their uses in antimicrobial chemotherapy are the milestone in the history of modern medicine and these agents are life-saving weapons against numerous infectious diseases. An antibiotic was a substance produced by one microorganism that selectively inhibits the growth or kills another microorganism. Antibiotics were initially viewed as ‘Wonder Drugs’ primarily because these were introduced at a time when only surgical drainage or spontaneous cures were available to treat serious bacterial infections” [2,4-6]. The mid-20th century was named the “antibiotic era”, and infectious diseases were believed to be eradicated by the end of the last century [1,6]. Contrary to the original belief, over a period of 50 years since the first antibiotic was approved for human use, many antibiotics lost efficacy due to safety concerns and the development and spread of antimicrobial resistance (AMR). Many microorganisms are continuously evolving and developing resistance and such resistant pathogens, ‘superbugs’, are rendering previously active antibiotics ineffective [1,5,6]. Much of the medical and pharmaceutical research and development has been focused on lifestyle disorders and cancer and many antibiotics were not approved since the 1980s entering the ‘post-antibiotic era’ wherein resources for discovery and development of active and effective antibiotics are limited.

1.1 Irrational Use of Antibiotics
It is highly essential to use antibiotics prophylactically in hospitalized patients for various surgeries, therapeutically in primary care settings for various community-acquired infections (CAIs), to prevent and treat infections in various invasive procedures and hospital acquired infections (HAIs) adhere to international, national and hospital antibiotic use policies [7-10]. Moreover, antibiotic use is compulsory in certain unavoidable clinical conditions, such as sepsis and in immunocompromised patients who are at risk of developing mixed infections” [9-12]. "Inarguably, when there are no clinical practice guidelines, there is a need and necessary to use antibiotics for prophylactic as well as therapeutic purposes
in the recent unprecedented pandemic time due to coronavirus disease (COVID-19) and in multisystem inflammatory syndrome in children (MIS-C) associated with SARS-CoV-2, the virus that causes COVID-19” [11,12]. However, inappropriate and irrational use, such as incorrect use, misuse, overuse, and often real abuse of antibiotics leads to the development of tolerance or antibiotic resistance from the time it is first employed [2,13]. Indeed, antibiotic consumption is most prevalent in many developing and developed countries. It is reported that India is among the highest use in the world and antibiotics sales continue to peak rapidly despite the prevalence of infectious diseases remaining stagnant [10,11,14]. It is a well-known fact that bacteria survive antibiotics challenges by selection pressure, undergo mutations, evolve as resistant, and regrow leading to the emergence and spread of new infectious diseases are one of the greatest life threats to human health. Consequently, the use of antibiotics is compromised, and several microorganisms, including bacteria are developing and spreading antibiotic resistance [2,14-16]. Therefore, AMR is increasing worldwide due to increased prescription, dispensing, over-the-counter sales, and consumption of antibiotics and is now considered as a threat to global health and sustainable economic development.

1.2 Antimicrobial Resistance

“AMR occurs when microbes, such as bacteria, viruses, fungi, and parasites undergo mutations and develop resistance that makes several previously effective medications ineffective and renders such infections untreatable. The term antibiotic resistance is a subset of AMR, as it applies to bacteria that become resistant to antibiotics. Resistant microbes are more difficult to treat, it requires higher doses, alternative medications, and multidrug treatment which may prove more toxic and may also be more expensive. In 2019 alone, an estimated 4.95 million deaths were associated with bacterial AMR which also implies the economic burden of the country” [16-18]. “Among the antibiotic-resistant bacteria, ‘ESKAPE’ pathogens, including, Enterococcus faecium, Staphylococcus aureus, Klebsiella pneumoniae, Acinetobacter baumannii, Pseudomonas aeruginosa, and, Enterobacter species, cause the majority of hospital infections with a higher rate of mortality” [19]. “In addition, other pathogens such as methicillin-resistant Staphylococcus aureus (MRSA), and vancomycin-resistant enterococci (VRE), effectively escape the effects of many antibacterial drugs which may lead to a longer hospital stay and decreasing workforce productivity” [20]. Moreover, most of the microbes exposed to one class of antimicrobials for a longer period develop AMR wherein cross-resistance becomingly troublesome. Microbes resistant to multiple antimicrobials (at least three different classes of antimicrobials) are called multidrug-resistant (MDR). Those bacteria that are considered extensively drug-resistant (XDR) or totally drug-resistant (TDR) are sometimes called ‘superbugs’ [8,9,21]. “There is a high risk of entering into a “post-antibiotic era”, a period in which bacteria have become resistant to existing antibiotics and the antibiotics no longer work” [13,22].

1.3 WHO Global Action Plan to Tackle Antimicrobial Resistance

“In order to tackle AMR, the World Health Organization (WHO) has developed Global Action Plan in 2015 keeping view of implementation in various nations across the world” [23]. “This plan is based on 5 objectives, including first, to improve awareness and understanding of AMR, second, to strengthen knowledge and generate a large amount of data, third, to reduce the incidence of infections through effective hygiene measures and good hygiene practice, fourth, to optimize the use of antimicrobial drugs in human and animal health by surveillance and stewardship programs, and fifth, to increase investment in new drugs, diagnostic tools, vaccines, and other interventions for discovery and development of new treatment modalities. In 2016, WHO listed the world’s leading antibiotic-resistant bacteria and priority pathogens, for which there is an utmost need for new antibacterial agents for effective treatments against superbugs, and MDR and XDR pathogens that are resistant to traditional antibiotics” [11,13,22,23]. “Since 2017, U.S. Food and Drug Administration (FDA) and the European Medical Agency (EMA) have approved several new antibiotics with predominant activity against Gram-negative bacteria, including plazomicin, eravacycline, cefiderocol, cefazidime/avibactam, cefotolozane/tazobactam, meropenem / vaborbactam (combination of beta-lactam with beta-lactamase inhibitors), imipenem-cilastatin / relebactam, and temocilllin, a beta-lactam antibiotic effective against Gram-negative bacteria” [1,6,24].
2. TETRACYCLINE ANTIBIOTICS

Tetracyclines have been one of the first antibiotics used to treat infections, they possess many properties considered ideal for antibiotic drugs, including a broad spectrum of activity against Gram-positive and Gram-negative pathogens, proven clinical safety, acceptable tolerability, low adverse effect profile, and availability of intravenous and oral formulations for most members of the class [25,26]. Tetracyclines are divided into three generations based on their spectrum of activity, and pharmacological properties such as half-life and binding to plasma protein (Table 1) [17,27].

2.1 Development of Tetracycline Antibiotics

Serendipitous discovery and extended use of earlier penicillins and streptomycin in the early 1940s have broadened the scope of identifying and developing newer antibiotics. With the advances in microbiology, biochemistry, and fermentation technology, several pharmaceutical scientists and academic researchers have focused on identifying several lesser-known microbes from soil samples collected across the globe and started isolating and discovering new antibiotics produced by such microorganisms. Owing to these efforts, aureomycin, the first tetracycline was discovered in 1945 and led to a series of tetracyclines development during the last century. Chlortetracycline is the synthetic form of aureomycin, a natural tetracycline antibiotic, which was discovered at Lederle Laboratories under the supervision of scientists Yellapragada Subbarow and Benjamin Minge Duggar [1,3,17]. Aureomycin was extracted from Streptomyces aureofaciens and was approved by the FDA in 1948 for medical use in humans. In the following years, terramycin, the second tetracycline, obtained from Streptomyces rimosus, and its synthetic form oxytetracycline were discovered and approved for human use in 1950 [3,17,25]. Molecular optimizations and structural changes in chlortetracycline lead to the discovery of tetracycline in 1953. Due to its favorable pharmacokinetic profile and enhanced water solubility, tetracycline is considered widely successful first-generation tetracyclines. The second-generation tetracyclines including methacycline, doxycycline, and minocycline were approved due to reduced risk of toxicity, better pharmacokinetic profiles, and extended antimicrobial spectrum were approved in the early 1970s (Table 1 and Table 2) [2,27,28].

Table 1. Classification of tetracyclines based on generations [13,27,28]

<table>
<thead>
<tr>
<th>Generation</th>
<th>Obtaining method</th>
<th>Agents</th>
</tr>
</thead>
<tbody>
<tr>
<td>First</td>
<td>Biosynthesis</td>
<td>Chlortetracycline, oxytetracycline, tetracycline, demeclocycline</td>
</tr>
<tr>
<td>Second</td>
<td>Semi-synthesis</td>
<td>Doxycycline, minocycline, lymecycline, meclocycline, methacycline, rolitetracycline</td>
</tr>
<tr>
<td>Third</td>
<td>Semi-synthesis</td>
<td>Tigecycline, omadacycline, sarecycline</td>
</tr>
<tr>
<td></td>
<td>Total synthesis</td>
<td>Eravacycline</td>
</tr>
</tbody>
</table>
2.3 Mechanism of Action of Tetracyclines

Tetracyclines inhibit protein synthesis by inhibiting the association of aminoacyl-tRNA with the bacterial ribosome and binding with high affinity to a 30s ribosomal subunit during translation. Then the penetration of aminoacyl-tRNA into the acceptor site (A) on the bacterial ribosome is blocked, which leads to the cessation in the incorporation of amino acid residues in the process of elongation of the polypeptide chain (Fig. 1). Thus, the bacteriostatic activity of antibiotics is achieved by stopping protein synthesis [28,29].

A few older tetracycline drugs are mostly used to treat specific microbes resistant to antibiotics. Similar to the classical tetracyclines, tigecycline, the first of the new generation tetracyclines, exhibits broad-spectrum antibacterial activity. Moreover, the three newer agents approved eravacycline, sarecycline, and omadacycline showed extended and unique antibacterial activity retaining the broad-spectrum activity of previous tetracyclines [17,27,28]. The antibacterial spectrum of newer agents comprises several resistant bacteria, including those resistant to older tetracyclines. Sarecycline is considered a narrow-spectrum tetracycline due to its highly selective antibacterial activity against *Cutinebacterium acnes* [30]. Exploratory research identified non-antibiotic properties of several tetracyclines, particularly the anti-inflammatory and neuroprotective effects of minocycline and sarecycline, possibly attributed partly to their inhibition of microglial activation and in part by modulating oxidative stress [31-33]. However, the exact mechanisms are not well elucidated and understood. Recently, the novel tetracyclines-exploiting strategy for neuroprotection based on their antiamyloidogenic, anti-inflammatory, antiapoptotic, and antioxidant activities has emerged for the treatment of Alzheimer’s and Parkinson’s diseases. Nevertheless, the challenges of repurposing tetracyclines are further hampered by safety issues, such as antibiotic-induced alteration of microbiota in the gut and consequent dysbiosis and the development and spread of AMR due to long-term antibiotic use.

![Fig. 1. Mechanism of action of tetracyclines](image)

*The tetracyclines reversibly bind to the 30S subunit of the bacterial ribosome thereby preventing the binding of acyl-transfer RNA (tRNA) to the ribosome. Thus, tetracyclines by inhibiting protein synthesis that stops the growth and replication of the bacterial organism leading to a bacteriostatic effect.*
3. THIRD-GENERATION TETRACYCLINE ANTIBIOTICS

3.1 Tigecycline

Tigecycline is a unique glycylcycline antibiotic approved by the US FDA in 2005. It has susceptibility to clinically important MDR nosocomial and community-acquired bacterial pathogens and also against a broad range of Gram-negative and Gram-positive bacteria species. It inhibits the translation elongation step by binding to the ribosome 30S subunit (five times stronger than other tetracyclines) and also prevents aminoacylated tRNA accumulation in the ribosomal A site even in the presence of ribosomal protecting efflux pumps [28,34]. MDR Gram-negative pathogens, such as A. baumannii and ESBL-producing K. pneumoniae and E. coli are highly susceptible to tigecycline. It is also active against VRE, MRSA, penicillin-resistant Streptococcus pneumoniae, anaerobes, and ‘atypical’ bacteria. However, it is not active against P. aeruginosa and Proteus, Morganella, and Providencia species [35]. Moreover, parenteral tigecycline was poorly tolerated and ineffective when tried as an agent in multidrug regimens for salvage therapy of Mycobacterium abscessus infection and an alternative option of oral formulation is not available. Therefore, omadacycline and eravacycline may represent a new therapeutic option for treating M. abscessus complex infections [36,37]. Recent reports show that tigecycline remained active even against Gram-positive and Gram-negative bacteria, at the same time resistant strains such as fluoroquinolone and broad spectrum β-lactam-resistant Enterobacteriaceae, vancomycin-resistant E. faecium, has increased during the same period [34,35]. Evidence exists that tigecycline is highly effective for the treatment of severe Clostridioides difficile infection in whom previous-generation tetracyclines are ineffective. In addition, Coxiella spp., Rickettsia spp., and MDR Neisseria gonorrhoea strains showed in vitro susceptibility to tigecycline, indicating its possible use in the treatment of such infections [38].

Tigecycline is indicated for the treatment of complicated skin and skin structure infections (SSSIs) caused by E. coli, E. faecalis (vancomycin-susceptible isolates), Staph. aureus (MSSA and MRSA isolates), S. agalactiae, S. anginosus group. (Include S. anginosus, S. intermedius, and S. constellatus), S. pyogenes, E. cloacae, K. pneumoniae, and B. fragilis. It is also indicated for the treatment of complicated intra-abdominal infections (cIAIs) caused by Citrobacter freundii, E. cloacae, E. coli, K. oxytoca, K. pneumoniae, E. faecalis (vancomycin-susceptible isolates), Staph. aureus (MSSA and MRSA isolates), S. anginosus group (includes S. anginosus, S. intermedius, and S. constellatus), B. fragilis, B. thetaiotaomicron, B. uniformis, B. vulgatus, Cl. perfringens, and Peptostreptococcus micros. Further, it is also approved for the treatment of community-acquired bacterial pneumonia (CABP) caused by S. pneumoniae (penicillin-susceptible isolates), including cases with concurrent bacteremia, H. influenzae (beta-lactamase negative isolates), and L. pneumophila [37,38]. It is administered as an intravenous infusion for a duration of about 30–60 min every 12 h. The recommended initial dose of tigecycline is 100 mg followed by 50 mg every 12 h and the duration of treatment with tigecycline for cSSTs or cIAIs and CAP is 5–14 and 7–14 days, respectively (Table 2) [39,40]. As the tetracyclines tigecycline also exhibits the same adverse events such as gastrointestinal symptoms commonly nausea, vomiting, anorexia, and other rare events including injection site irritation, pain, and swelling [41,42]. It is to be noted that tigecycline has no approved indication for the treatment of diabetic foot infection or for hospital-acquired or ventilator-associated pneumonia, in which a high rate of mortality is reported [39]. The US FDA has already warned that there is an increased risk of death resulting from complications of infection, worsening infections, or other underlying medical conditions when intravenous tigecycline is used for approved and non-approved uses. Presently, tigecycline is a reserve antibiotic for use in situations when alternative antibiotic therapies are not appropriate [39,41,43]. Importantly, advice and concordance between health care professionals, patients, and their caregivers are highly essential for using tigecycline in such situations.

3.2 Omadaclycline

Omadaclycline, a novel aminomethyl tetracycline antibiotic, has been developed to combat the AMR resistance to earlier tetracyclines [43]. It possesses excellent activity against many bacterial species and reversibly binds to the 30S ribosomal subunit and inhibits protein synthesis. Due to its reversible binding to the microbial ribosome, it acts as a bacteriostatic. In vitro omadaclycline has demonstrated bactericidal activity against Haemophilus influenzae, Streptococcus pneumoniae, and Moraxella catarrhalis [44]. Mycobacterium abscessus complex (MÁBC)- non-tuberculous mycobacteria
that include (*M. abscessus*, *M. bolletii*, and *M. massiliense*) which are the major cause of human pulmonary or skin and skin structure infections (SSSI) has acquired resistance to older tetracyclines but now sensitive to omadacycline [45-47]. The characteristics of omadacycline include broad-spectrum activity, and overcoming the two primary mechanisms of tetracycline resistance, efflux, and ribosome protection. It has shown efficacy and is well-tolerated when used for acute bacterial SSTIs and CAP [17,27,48]. The other two pharmacological benefits of its effective oral administration include good oral bioavailability and lack of glycycline-induced dose-limiting nausea and vomiting [43]. Omadacycline has minimal known drug-drug interactions and should be administered in a fasting state, avoiding dairy and cation-containing products for at least 4 h after dosing [48]. The chemical structure of omadacycline is similar to tigecycline, and it is the derivative of minocycline [26,27]. With increasing awareness and surveillance of antibiotic utilization, antimicrobial stewardship programs have continuously been considering utilization of omadacycline as potential therapeutic option in the treatment of infections caused by MSSA isolates as well as antibiotic-resistant and MDR Gram-positive bacteria, including MRSA, VRE, including vancomycin-resistant *E. faecium* and *E. faecalis*, penicillin and tetracycline-resistant *S. pneumoniae* and *S. viridans*, erythromycin-resistant *S. agalactiae*. Moreover, omadacycline is also equally effective against Gram-negative pathogens, such as *E. coli*, *K. pneumoniae*, *P. aeruginosa*, *E. cloacae*, *H. influenzae*, *H. parainfluenzae* Citrobacter spp., and *P. mirabilis*. Further, its antibacterial spectrum is also extended to respective ESBL and carbapenem-resistant *Enterobacteriaceae* (CRE) phenotypes as well as ceftazidime-resistant strains [26,46]. It also exhibits potent *in vitro* activity against non-mycobacterial atypical organisms, including *Mycoplasma pneumoniae* and *M. hominis*, *Legionella pneumophila*, and *Chlamydia pneumoniae*. Omadacycline could positively affect hospitalization-associated expenses for both bacterial CAP and acute bacterial SSTIs by providing another oral agent active against resistant Gram-positive pathogens [48].

It is indicated for the treatment of adult patients with community-acquired bacterial pneumonia (CABP) based on susceptible patterns against several microorganisms. Further, it is also indicated for the treatment of adult patients with acute bacterial skin and skin structure infections (ABSSSI) caused by several resistant bacteria, including *Staph. lugdunensis*, *S. pyogenes*, and *S. anginosus* group. (Includes *S. anginosus*, *S. intermedius*, and *S. constellatus*) [48,49]. It is indicated for CABP given once-daily oral and intravenous administration [48,49]. For injection, omadacycline is available as single-dose vials containing 100 mg of omadacycline as a sterile lyophilized powder. It is recommended intravenous dose of 200 mg should be administered slowly as an infusion over 60 min duration whereas a 100 mg dose is administered over 30 min (Table 2) [44,49]. Besides, omadacycline is an ideal choice even in outpatient and ambulatory care settings wherein such treatment options enable healthcare facilities to discharge patients who cannot take other oral antibiotics and reduce the duration of hospital stay. Adding to this, such a therapeutic option would be a preferable choice for treatment on an outpatient basis which could further reduce the cost and risk of hospitalization(s) [50]. Overall, omadacycline is well tolerated, although nausea and vomiting are frequently reported [49]. Notwithstanding this favorable safety profile, omadacycline caused a moderate increase in heart rate during the treatment period and such effects are possibly mediated through muscarinic m2 receptors [49,51]. On the other hand, omadacycline was reported to cause an increase in hepatic enzymes, albeit relatively low. Indeed, it shares many of the undesirable side effects of tetracyclines, such as tooth discoloration, inhibition of bone growth, etc., [49].

### 3.3 Eravacycline

Eravacycline is a novel broad-spectrum tetracycline that has been developed to overcome tetracycline resistance. It is indicated for the treatment of intra-abdominal infections caused by susceptible microorganisms and has a broad antibacterial spectrum that includes all common clinical pathogens except *P. aeruginosa*, including Gram-negative and Gram-positive aerobic and anaerobic strains [52]. It is a totally synthetic fluoroquinolone specifically designed to overcome tetracycline-acquired resistance associated with ribosomal protection mechanisms and efflux pumps [53]. It also shows
Table 2. Antibacterial spectrum, pharmacokinetics, dosage regimen, safety, and indications of newer tetracyclines [17,26,35,53,68]

<table>
<thead>
<tr>
<th>Drug name</th>
<th>Spectrum</th>
<th>Pharmacokinetics ($t_{1/2}$, $T_{max}$, $C_{max}$, MIC)</th>
<th>Dose and duration</th>
<th>Side effects</th>
<th>Indications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tigecycline</td>
<td>Broad spectrum</td>
<td>$t_{1/2}$: 1.05-2.34 h $C_{max}$: 0.42-11.1 µg/mL $T_{max}$: 2.0-2.8 h $MIC$: ≤ 2 µg/mL</td>
<td>An initial dose of 100 mg followed by 50 mg every 12 h for 5–14 and 7–14 days</td>
<td>Nausea, vomiting, injection site irritation, pain, swelling, and anorexia</td>
<td>cSSTIs, cIAIs (cholecystitis, gangrenous perforated diverticulitis, appendicitis, peritonitis) CAP</td>
</tr>
<tr>
<td>Omadacycline</td>
<td>Gram-positive and negative aerobic pathogens</td>
<td>$t_{1/2}$: 16-17 h $T_{max}$: 2.5 h $C_{max}$: 0.3-4.5 µg/mL $MIC$: 0.06 and 0.125 µg/mL</td>
<td>An intravenous dose of 200 mg infusion over 60 min, 100 mg dose is administered over 30 min</td>
<td>Nausea, vomiting, diarrhea, tooth discoloration, inhibition of bone growth, increased heart rate, increased hepatic enzymes</td>
<td>cSSTIs CAP UTIs</td>
</tr>
<tr>
<td>Eravacycline</td>
<td>Broad-spectrum against aerobic and anaerobic Gram-negative and positive bacteria, except P. aeruginosa and Burkholderia cenocepacia</td>
<td>$t_{1/2}$: 20 h $C_{max}$: 1 h $T_{max}$: 1.5-2 h $MIC$: 0.5-2 µg/mL</td>
<td>1 mg/kg every 12 h, i.v. infusion approximately over 60 min every 12 h for 4 to 14 days</td>
<td>ISRs, nausea, vomiting, diarrhea hypotension, and wound dehiscence</td>
<td>cIAIs (diverticulitis, appendicitis, intra-abdominal abscess, cholecystitis, gastric and duodenal perforation, intestinal perforation, and peritonitis)</td>
</tr>
<tr>
<td>Sarecycline</td>
<td>Narrow-spectrum activity against aerobic and anaerobic Gram-negative bacteria, limited activity against Gram-positive organisms</td>
<td>$t_{1/2}$: 21-22 h $T_{max}$: 1.5-2.0 h $C_{max}$: 1.5-2.0 h $MIC$: 0.5 µg/mL</td>
<td>0.75, 1.5, 3.0 mg/kg, OD, for 12 weeks 33 to 54 kg: 60 mg 55 to 84 kg: 100 mg 85 to 136 kg: 150 mg No dose adjustments for hepatic/renal impairment</td>
<td>Nausea, vomiting, abdominal pain and discomfort, nasopharyngitis, sunburn, vulvovaginal candidiasis, and vulvovaginal mycotic infection</td>
<td>Acne vulgaris (inflammatory skin lesions of non-nodular with moderate-to-severe acne vulgaris in patients who are 9 years old and above)</td>
</tr>
</tbody>
</table>

CAP: Community-acquired pneumonia; cIAIs: Complicated intra-abdominal infections; cSSTIs: Complicated skin and soft tissue infections; MIC: Minimum inhibitory concentration; UTI: Urinary tract infection
good antibacterial activity against MDR bacteria, including Enterobacteriaceae and A. baumannii, expressing extended-spectrum. It is more effective than omadacycline against Gram-negative and broad-spectrum beta-lactamase-producing bacteria [53]. It is particularly active against all rapidly growing Mycobacterium species which includes M. abscessus subsp. abscessus, M. abscessus subsp. massiliense, M. chelonae, M. immunogenum, M. fortuitum, and M. mucogenicum groups [54]. It shares the same mechanism of action as other tetracyclines and the main effects are bacteriostatic and reversible, with some bactericidal activity against certain bacterial species. It was previously believed that eravacycline could be utilized in complicated urinary tract infections (cUTI), but failed to show efficacy in comparison to carbapenems. Despite this, it is characterized by a broad-spectrum antimicrobial spectrum with coverage of several resistant strains, such as MRSA, VRE, CRE, tetracycline-resistant bacteria, as well as A. baumannii [55]. It has extended spectrum of antimicrobial activity against E. coli, K. pneumoniae, Citrobacter freundii, E. cloacae, Streptococcus anginosus group, Cl. perfringens, Bacteroides species, and Parabacteroides distasonis, Klebsiella oxytoca, Enterococcus faecalis, Enterococcus faecium, Staph. aureus. However, it is not indicated for the treatment of cUTI.

It is indicated for the treatment of complicated intra-abdominal infections (cIAI) caused by E. coli, K. pneumoniae, Citrobacter freundii, E. cloacae, K. oxytoca, E. faecalis, E. faecium, Staph. aureus, S. anginosus group, Cl. perfringens, Bacteroides species, and Parabacteroides distasonis in patients 18 years or older [56]. In practice, it should be used only to treat or prevent infections that are suspected to be caused by susceptible bacteria. It has an advantage over other new tetracyclines as both oral and intravenous formulations are available. The recommended dosage regimen is 1 mg/kg every 12 h for slow intravenous infusion over 60 min every 12 h. The commonly reported adverse effects are infusion site reactions, nausea, vomiting, diarrhea, hypotension, and wound dehiscence. Though well studied for antibacterial activity against infections in adults, its safety and effectiveness in pediatrics have not been established. Moreover, dosage adjustment is warranted in patients with severe hepatic and/or renal impairment (Table 2) [57,58]. To some extent, resistance in some bacteria to eravacycline is associated with up-regulated, non-specific intrinsic MDR efflux, and target-site modifications, such as 16s rRNA or certain 30S ribosomal proteins [53]. It is approved only for the treatment of CAP and acute SSTIs, such as appendicitis, cholecystitis, diverticulitis, gastric and duodenal perforation, intra-abdominal abscess, intestinal perforation, and peritonitis. Recent clinical trials also demonstrated its use in pylonephritis and cystitis. Though it has very less drug-drug interactions, it should be avoided with dairy and divalent cation-containing products for at least 4 h of duration after the dosing [52,56,58].

3.4 Sarecycline

Sarecycline is a novel, narrow-spectrum, once-daily, and oral tetracycline-class antibiotic approved in October 2018. It has potent activity against Gram-positive bacteria, including activity against multiple strains of Cutibacterium acnes, an anaerobic Gram-positive bacterium that causes acne lesions and possesses anti-inflammatory properties similar to other tetracyclines. Most of the older tetracyclines have broad-spectrum antibacterial activity, contrary to that sarecycline is less potent due to its activity against enteric aerobic Gram-negative bacteria and anaerobic bacteria is minimal [30,59-61]. Owing to its narrow-spectrum antibacterial activity, sarecycline is associated lower risk of potential adverse effects, thus making it a potential therapeutic choice for definitive treatment among antibiotics, including tetracyclines. In addition to this, sarecycline had shown low susceptibility to resistance over other tetracyclines. Importantly, it is active against erythromycin- and clindamycin-resistant C. acnes strains as well as tetracycline-resistant Staph. aureus [59,64]. It is well known that acne vulgaris affects almost everyone, particularly during teenage and young adult years, though over 40% of individuals still suffer from acne in adulthood as well. Keeping in view of this, sustained efforts have been made to identify the safest, well-tolerated, and most effective treatments. [62,63,65]. Despite many studies being conducted to determine the antimicrobial spectrum of sarecycline compared to other tetracyclines, it is still a narrow-spectrum antibiotic with the purpose to reduce and tackle AMR [30,64]. It exerts its antimicrobial effect mainly as a ribosomal protein synthesis inhibitor. Intriguingly, it has a unique mechanism of action due to it has the longest and largest chemical moiety attached at the carbon-7 (C7) position of ring D of the four-ring core thus exerting
antibiotic effect by binding to the decoding center of the 30S subunit of the bacterial ribosome, thereby inhibiting mRNA to protein translation [29,30,66]. It is active against both methicillin-susceptible (MSSA) and methicillin-resistant (MRSA) strains of S. aureus as well as S. epidermidis, but less active than doxycycline and minocycline by twofold. Indeed, it is more active than tetracycline and doxycycline against S. pyogenes, S. agalactiae, E. faecalis, and E. faecium (both vancomycin susceptible and resistant).

It is used as a narrow-spectrum antibiotic specifically indicated for the treatment of inflammatory lesions of non-nodular moderate-to-severe acne in patients 9 years old and above [62,63,67]. The treatment regimen for acne includes one sarecycline tablet per day (equivalent to 1.5mg/kg/day) administered orally as 60 mg, 100 mg, or 150 mg with or without food for a duration of up to 40 weeks (Table 2) [60,67,68]. Patient education and medication adherence are highly essential while taking sarecycline. It is recommended that patients should ingest the tablet at the same time each day at least one hour before or two hours after eating for beneficial and desirable therapeutic outcomes. It is recommended not to use this drug beyond 12 months since its safety beyond 12 months has not been established yet [67]. The common adverse effects of sarecycline include nausea, bloody stools, stomach irritation, phototoxic adverse effects, light-headedness, vertigo, dizziness, and abnormal pressure in the brain. Apart from the threat of developing AMR, the use of some of oral antibiotics is often limited by their vestibular side effects, such as dizziness and vertigo. It is reported that sarecycline poorly crosses the blood-brain barrier due to its weak lipophilicity when compared with other tetracyclines which may explain relatively lower rates of vestibular-related side effects observed in clinical trials [69]. Animal toxicity studies, it has shown skeletal defects in offspring, decreased fertility, and decreased spermatogenesis. Therefore, it is contraindicated in pregnant and breastfeeding women due to the risk of teratogenic effects [67].

4. CONCLUSIONS

The emergence and tackling of antimicrobial resistance is a crisis for the health and wealth of nations across the world. Irrational and inappropriate use of antibiotics is one the main cause of the development and spread of AMR. Owing to this, most of microorganisms are evolving due to selection pressure, developing resistance, and rendering antibiotics ineffective in treating many infectious diseases. The discovery and emergence of newer tetracyclines have shown remarkable effects against bacteria that are resistant to previous antibiotics including older tetracyclines. Recently approved tetracyclines, omadacycline, eravacycline, and sarecycline as well as tigecycline, have the advantage of a superior potency against both Gram-positive and Gram-negative aerobic as well as anaerobic MDR bacteria. These drugs also have a broad spectrum of activity which is very advantageous in treating many infectious diseases. These newer tetracyclines act like magic bullets, particularly against antibiotic-resistant pathogens and 'superbugs', and had shown promising results in treating infections caused by antibiotic-resistant, MDR, and XDR bacteria. Through these new antibiotics categorized as the third-generation tetracyclines, their discovery, development, and subsequent approval with specified indications are important milestones in medicine, particularly, to rationalize appropriate use and to minimize the development and spread of AMR. Essentially, these agents serve as leading molecules and maneuver for the discovery and development of newer antibiotics in the post-antibiotic era. Inarguably, continuous surveillance on antibiotic utilization and implementation of antibiotic stewardship programs are warranted to help select appropriate treatment, to optimize therapeutic outcomes, minimize AMR, and sustain therapeutic efficacy for the future.

COMPETING INTERESTS

The authors have declared that no competing interests exist.

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