Posaconazole in the management of refractory invasive fungal infections

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Abstract: The rising incidence of invasive fungal infections due to the expanding population of immunocompromised hosts and the increasing prevalence of fungal resistance has led to the need for novel antifungal agents. Posaconazole, a new member of the triazole class has demonstrated in vitro activity against a broad spectrum of fungi and clinical activity against various fungal pathogens, including Aspergillus spp., Candida spp., zygomycetes, and Fusarium spp. To date, posaconazole has been approved for prophylaxis of invasive fungal infections in stem cell transplant recipients with acute graft versus host disease (GVHD) and neutropenic patients receiving intensive induction chemotherapy for acute myelogenous leukemia and myelodysplastic syndrome. In addition, it has been licensed for use in oropharyngeal candidiasis and for salvage therapy in invasive aspergillosis, fusariosis, coccidioidomycosis, chromoblastomycosis, and mycetoma. Posaconazole is the only azole with activity against zygomycetes and other difficult-to-treat fungi, representing a potential treatment option for refractory invasive mycosis. This article reviews available preclinical and clinical data of posaconazole, focusing on its role in the treatment of refractory invasive fungal infections.

Keywords: posaconazole, refractory invasive fungal infections, salvage therapy

Introduction

Invasive fungal infections (IFI) have increased significantly as a leading cause of life-threatening conditions in immunocompromised patients over the past two decades (Marr et al 2002; Wisplinghoff et al 2004). This is primarily due to the rise of at-risk individuals comprising immunocompromised patients with prolonged neutropenia or advanced HIV infection, and patients who have undergone allogeneic hematopoietic stem cell transplantation (HSCT) or solid organ transplantation. Improvement in anticancer treatment, greater duration and intensity of immunosuppression, and the variety of available antimicrobial therapies have influenced the spectrum of pathogens associated with IFI. Although Candida and Aspergillus spp. remain the principal causes of most IFI, mycoses due to more unusual fungal pathogens like Cryptococcus, Coccidioides, Histoplasma, and agents of zygomycosis (primarily species of Rhizopus, Mucor, Cunninghamella, Apophysomyces, Absidia, and Rhizomucor) have become more prevalent in recent years. In fact, the effectiveness of routine fluconazole prophylaxis has resulted in a reduction of C. albicans infections, but has also caused a shift to non-albicans Candida spp., ie, C. krusei and C. glabrata – representing fluconazole-resistant strains – and C. parapsilosis (Pfaller et al 1998; Marr et al 2000; Baran et al 2001). To date only four classes of antifungal agents have been approved for the treatment of IFI: the azoles such as itraconazole, ketoconazole, fluconazole, and voriconazole; the polyenes, the most prominent of which is amphotericin B; fluconosine, a drug inhibiting fungal RNA and DNA synthesis; and the echinocandins, such as caspofungin, anidulafungin, and micafungin (Table 1). Failure rates of these agents, including the recently introduced agents, such as voriconazole and caspofungin, are
## Table 1 Drug Profiles

| Antifungal class | Polyene | Lipid formulations |
|------------------|---------|--------------------|
| Amphotericin B   | intravenous | intravenous |
| Echinocandins    | intravenous | intravenous |
| Fluconazole      | intravenous | oral |
| Itraconazole     | intravenous | oral |
| Voriconazole     | oral |
| Posaconazole     | oral |

### Mechanism of Action
- **Polyene**: Interaction with ergosterol, intercalation of fungal membrane, increased membrane permeability
- **Azoles**: Inhibition of cytochrome P450 14 alpha-demethylase, accumulation of lanosterol leading to perturbation of fungal cell membrane, fungistasis
- **Echinocandins**: Inhibition of cell-wall glucan synthesis leading to susceptibility of fungal cell to osmotic lysis

| Loading dose/day | Usual maintenance dose/day | Frequency of dosing | Half-life (hours) | Oral bioavailability | CNS penetration | Route of elimination | Dose reduction in renal failure | Dose reduction in hepatic failure | Major toxic effects |
|------------------|---------------------------|---------------------|------------------|----------------------|-----------------|----------------------|-----------------------------|------------------------|------------------|
| No               | 0.6–1.5 mg/kg daily       | daily               | 24               | NA                   | poor            | slow renal excretion | consider lipid formulation | data not available | dose-limiting renal toxicity, fever, chills |
| No               | 400-800 mg iv/po daily    | daily               | 30               | >90%                 | data not available | renal               | not required               | data not available | less toxic than amphotericin B |
| No               | 400 mg iv / 4 days       | twice daily         | 56–64            | 55%                  | very good       | fecal (primarily) renal | not required               | data not available | well tolerated occasional GI toxicity |
| No               | 400 mg po /3 days        | twice daily         | dose dependent   | 96%                  | moderate        | focal renal          | not required               | data not available | liver and GI toxicity |
| No               | 12 mg/kg iv               | 2–4 divided doses  | 20               | variable*            | moderate        | focal renal          | not required               | not required         | drug interactions, visual, liver, skin toxicity GI and liver intolerance |
| No               | 600–800 mg                |                      |                  |                      | moderate        | focal renal          | not required               | required               | well tolerated occasional histamine mediated symptoms |
| No               |                          |                      |                  |                      | moderate        | focal renal          | not required               | required               | GI intolerance |
| No               |                          |                      |                  |                      | very good       | renal               | not required               | not required         | marrow suppression |

### Abbreviations
- NA, not applicable; iv, intravenous; po, orally; CNS, central nervous system; GI, gastrointestinal.
- *enhanced with food (2-fold), with high fat meal (4-fold)
still high – up to 60%–80% in allogeneic HSCT recipients (Barnes and Marr 2007). Hence there is an obvious need for new, more potent, broad-spectrum antifungal agents.

Posaconazole (Noxafil®; Schering-Plough), an extended-spectrum triazole has been recently approved for prophylaxis and treatment of refractory IFI by the EMEA and the FDA. The aim of this review is to summarize the in vitro and clinical data available on posaconazole with special focus on refractory invasive mycoses.

Mode of action and mechanism of resistance

Posaconazole is a lipophilic triazole with structural similarity to itraconazole. Theazole class of drugs inhibits the cytochrome P450 (CYP450)-dependent lanosterol 14 alpha-demethylase (CYP51), an enzyme required for ergosterol production which is an essential sterol component in the cell membrane of fungal pathogens but is not present in mammalian cells (Munayyer et al 2004; Wexler et al 2004). Exposure to posaconazole is a more potent inhibitor of sterol synthesis in Aspergillus fumigatus and A. flavus than either itraconazole or voriconazole. Posaconazole has an exceptional high affinity to CYP51. Since posaconazole has a chemical structure different from that of fluconazole and voriconazole, it can interact with an additional domain of the target so that it inhibits even mutated strains resistant to fluconazole and voriconazole (Munayyer et al 2004; Hof 2006). CYP51 is encoded by the ERG11 gene in Candida albicans and is present in almost all yeasts and molds, with the exception of Pneumocystis spp. and Phytium spp. (Hof 2006). Inhibition of CYP51 results in depletion of ergosterol from the fungal cell membrane and accumulation of methylated sterol precursors. This results in membrane instability, increased permeability, and inhibition of cell growth (Kwon et al 2007).

Mechanisms of resistance toazole antifungals have been reported to occur via both mutation of CYP51 and upregulation of genes controlling drug efflux pumps (Lupetti et al 2002; Hof 2006). The target site for the azoles is the ERG11 gene product encoding the cytochrome P450 lanosterol 14 alpha-demethylase, an essential enzyme in the biosynthetic pathway of ergosterol. Many of the identified CYP51 mutations map to the active site of the enzyme, thereby reducing the binding affinity of the respective azoles to their cellular target. Mutations seem to preferentially affect binding of fluconazole and voriconazole versus itraconazole and posaconazole, probably because of the long side chain of the latter two agents (Xiao et al 2004; Kwon and Mylonakis 2007). Based on 3D models of CYP51 bound to azoles, these side chains occupy a specific channel within CYP51, and this additional interaction serves to stabilize the binding of these azoles to the mutated CYP51 proteins (Xiao et al 2004). The model also predicts that mutations that were previously shown to specifically impact posaconazole susceptibility in A. fumigatus and C. albicans act by interfering with the binding of the long side chain. Azoles passively diffuse into fungal cells and resistance of yeasts as well as molds can develop when transmembrane efflux pumps are activated, thereby decreasing the intracellular drug concentration. The transporters of the ABC family CDR1 and CDR2 together with the major facilitator efflux gene MDR1 have all been implicated in the development of fungal resistance to azoles. However, these efflux pumps seem to work less efficient when exporting posaconazole in comparison to the other azoles (Chau et al 2004; Akins 2005). CDR1 and CDR2 have been shown to efficiently transport fluconazole, ketoconazole, itraconazole, and voriconazole, whereas posaconazole is transported to a much lesser degree. MDR1 can specifically transport fluconazole but has no effect on posaconazole (Akins 2005).

Pharmacokinetics and metabolism

Posaconazole is available as an oral suspension administered at 800 mg daily in 2 or 4 daily doses for salvage treatment or 600 mg daily in 3 daily doses for prophylaxis. The pharmacokinetics of posaconazole have been studied in single- and multiple-dose studies in healthy volunteers. Collectively, these studies have established that posaconazole displays a linear, dose-proportional pharmacokinetic up to single doses of 800 mg daily (Courtney et al 2003, 2004; Krieter et al 2004). Posaconazole is orally bioavailable, with maximum concentrations reached approximately 10 hours post dose. The absorption of posaconazole is linear through the clinically useful dose of 400 mg every 12 hours; absorption is saturated at higher doses and, thus, loading doses are not possible (Krieter et al 2004). Likeitraconazole, food greatly affects posaconazole absorption. Absorption of the oral suspension increases approximately 2-fold when taken with food, and 4-fold with a high-fat meal (Courtney et al 2004). Multiple-dose studies demonstrated that splitting the dose increased the total amount of drug absorbed (Courtney et al 2003). Posaconazole has a large volume of distribution (486 L) – compatible with good tissue penetration – and is >95% protein bound. It undergoes limited hepatic metabolism, via UDP glucuronidation and less through the oxidative pathways of the CYP450 system. Elimination
occurs predominantly through fecal excretion (77%) and to a lesser extent through urinary excretion (14%) (Ghosal et al 2004; Krieter et al 2004). As a result of posaconazole’s high protein binding and extensive distribution, the terminal plasma elimination half-life is approximately 20 hours, and steady-state concentrations are reached within approximately 7–10 days after initiation of therapy (Krieter et al 2004). This might affect its use in primary therapy for IFI. In a small study comprising patients with varying degrees of hepatic dysfunction, a trend of increasing half-life indicating prolonged elimination was visible; however, the area under the curve (AUC) did not significantly change and therefore, at this time, no dose adjustment is suggested for patients with hepatic impairment (Courtney et al 2000). In a small study of 24 patients with mild to moderate renal impairment, there was no correlation between pharmacokinetics of posaconazole and mild (creatinine clearance 50–80 mL/min) to moderate (creatinine clearance 20–49 mL/min) renal dysfunction after a single 400 mg oral dose. The drug was not removed by hemodialysis. These data indicate that posaconazole can be administered to subjects with varying degrees of renal impairment without dose adjustment, and supplemental doses are not needed after hemodialysis (Courtney et al 2005; Ramsewak et al 2005). Mucositis in neutropenic stem cell transplant recipients appears to reduce posaconazole absorption but did not significantly affect mean total posaconazole exposure at steady state condition. Moreover, this reduction could be overcome by increasing the total dose and dosing frequency (Gubbins et al 2006).

**Drug interactions**

Drug interactions mediated by various CYP450 are common with the currently available triazole antifungals; however, recent results suggest that posaconazole may have an improved drug interaction profile compared with other triazoles (Wexler et al 2004). Unlike other azoles, posaconazole is not a significant substrate for the CYP450 enzymes and has been shown to inhibit only CYP3A4. Therefore, posaconazole has the potential to interact with other drugs that are metabolized through the 3A4 enzyme system (Ramsewak et al 2005). Pharmacokinetic studies in special populations revealed no necessity for dosage adjustment based on differences in age, gender, race, or renal or hepatic function (Groll and Walsh 2005). The effect of posaconazole on the pharmacokinetics of tacrolimus and ciclosporin was evaluated in healthy subjects and transplant patients revealing that clearance of these immuno-suppressive agents was significantly decreased while the half-life increased. Therefore, when used in combination with mold-active azoles, calcineurin inhibitor doses should be reduced by at least 50% and their blood or serum concentrations should be closely monitored (Groll and Walsh 2005; Gubbins 2007). The clearance of posaconazole increased 2-fold in the presence of rifabutin, a potent inducer of CYP3A4. Therefore, co-administration of these two drugs is not advocated. Likewise, concomitant use of posaconazole with phenytoin or cimetidine should be avoided because these drugs decrease posaconazole concentrations by approximately 50% and 40%, respectively (Torres et al 2005). No dose adjustments are needed when posaconazole is coadministered with drugs such as glipizide, zidovudine, or lamivudine (Groll and Walsh 2005). Co-administration with CYP3A4-metabolised statins (simvastatin, lovastatin, atorvastatin) is contra-indicated as is co-administration with non-sedating antihistamines, cisapride, quinidine, or halofantrine, which may induce QT prolongation and torsades de pointes (Metcalf and Dockrell 2007; Zonios and Bennett 2008). Because interactions caused by itraconazole have been extensively studied and both itraconazole and posaconazole inhibit CYP3A4, drugs with known interaction to itraconazole should be used with caution in posaconazole-treated patients (Zonios and Bennett 2008). Since most drug-drug interaction investigations involving posaconazole have not yet been published, at this time the best sources for potential interactions with posaconazole is the prescribing information.

**In vitro studies**

*Aspergillus* spp./activity against molds

Filamentous fungal pathogens are increasingly becoming a major cause of IFI in immunocompromised patients. Most of these infections are caused by *A. fumigatus* (90%), followed by *A. flavus, A. niger,* and *A. terreus* (Kwon and Mylonakis 2007). Voriconazole is generally regarded as treatment of choice for invasive aspergillosis (Herbrecht et al 2002; Walsh et al 2008). However, Pfaller evaluated the in vitro antifungal activities within the SENTRY Antimicrobial Surveillance Program in the US and Canada of the new triazole antifungal agents, including posaconazole, voriconazole, and itraconazole, as well as amphotericin B, against 239 clinical isolates of filamentous fungi. Overall, posaconazole was the most active compound, inhibiting 94% of isolates at a minimal inhibitory concentration (MIC) of ≤1 μg/mL, followed by voriconazole (91%), amphotericin B (89%), ravuconazole (88%), and itraconazole (70%). Posaconazole also exhibited excellent activity (MICs, 0.03–1 μg/mL)
against less common filamentous fungi tested such as *A. niger*, *A. flavus*, *A. versicolor*, and *A. terreus*, which is often resistant to amphotericin B. None of the triazoles were active against *Fusarium* spp. These data have been largely confirmed by a more recent report of the SENTRY study group (Diekema et al. 2003). Also, in studies assessing the activity of posaconazole against approximately 19,000 clinically important strains of yeasts and molds including 1,423 *Aspergillus* spp., isolates collected from 200 medical centers worldwide over a 10-year time span, posaconazole was more or equally active than the comparator drugs itraconazole, fluconazole, voriconazole, and amphotericin B in almost all molds tested. Remarkably, posaconazole was active against isolates of *Candida* and *Aspergillus* spp. that exhibited resistance to fluconazole, voriconazole, and amphotericin B and was much more active than the other triazoles against zygomycetes (Sabatelli et al. 2006). However, *A. niger* seems to be less susceptible to posaconazole (Espinel-Ingroff 2003). The clinical implications of raised posaconazole MICs are unclear since it may be possible to achieve clinical serum posaconazole concentrations higher than these MICs (Torres et al. 2005).

**Zygomycosis**

Zygomycosis is an increasingly emerging life-threatening infection that particularly affects patients with diabetes or malignancy. Posaconazole appears to be the only azole that demonstrates activity against most Zygomycetes. The most common pathogens of human disease are *Rhizopus* spp. (47%) and *Mucor* spp. (18%), followed by *Cunninghamhamella bertholletiae* (7%) and *Apophysomyces elegans* (6%) (Roden et al. 2005). Amphotericin B remains first line therapy, but posaconazole has shown in vitro activity against many strains of zygomycetes (Dannaoui et al. 2003; Sabatelli et al. 2006; Sun et al. 2002). A recent comparative study comprising 19,000 yeast and mold isolates, 86 of which were *Zygomycetes* with voriconazole and itraconazole showed that posaconazole exhibited the lowest MICs against *Zygomycetes* isolates (Sabatelli et al. 2006). In a study of 37 strains of *Zygomycetes*, posaconazole was effective against *Mucor* spp., *Rhizopus* spp., *Absidia* spp., *Rhizomucor* spp., *C. bertholletiae*, and *A. elegans* (Sun et al. 2002; Kwon and Mylonakis 2007).

**Candida** spp./activity against yeasts

The incidence of invasive candidiasis is increasing and *Candida* spp. are now the fourth most common cause of bloodstream infection (Gudlaugsson et al. 2003; Pappas 2006). The mortality attributable to invasive candidiasis ranges from 30% to 71% with significant variation between different *Candida* spp. (ie, in hematologic malignancies the mortality is highest for *C. albicans* and *C. tropicalis* and lowest for *C. parapsilosis*), as well as underlying conditions, comorbidity and therapeutic procedure (for example HSCT, corticosteroid therapy, antibiotics) (Pagano et al. 2006; Staber et al. 2007). *C. albicans* is the most common cause of candidemia in adults and children, and is responsible for 40%–77% of cases (Pappas et al. 2006). The in vitro activity of posaconazole has been tested against over 10,000 clinical *Candida* isolates (Pfaller et al. 2004; Cuenca-Estrella et al. 2006; Sabatelli et al. 2006). In vitro, posaconazole is highly active against *Candida* spp. The drug is more active than itraconazole and fluconazole against all *Candida* spp. and *Cryptococcus neoformans*. During the ARTEMIS global antifungal surveillance program, 4,169 clinical isolates of *Candida* spp. were investigated for in vitro susceptibilities of voriconazole, posaconazole, and fluconazole (Pfaller et al. 2004). Both voriconazole and posaconazole were more active than fluconazole against all *Candida* spp. and *C. neoformans*. Posaconazole was the most active azole tested, and was unaffected by fluconazole resistance in *C. albicans* isolates and also had activity against *Candida* spp. that are commonly azole resistant, including *C. glabrata* and *C. krusei*. Posaconazole exhibited fungistatic and fungicidal activity in vitro and in vivo for most *Candida* spp. isolates and inhibited 97% of *Candida* spp. isolates at concentrations of 1 μg/mL or below (Pfaller et al. 2001; Espinel-Ingroff 2003). *C. albicans* is the most susceptible species of *Candida* whereas *C. glabrata* and *C. pelliculosa* are the least susceptible (Pfaller et al. 2001, 2004; Carrillo-Munoz et al. 2005). Nevertheless, posaconazole showed good activity against fluconazole resistant strains especially Candida glabrata, although it was slightly less active than voriconazole (Sabatelli et al. 2006; Pfaller et al. 2008).

**Cryptococcus neoformans**

Cryptococcosis is one of the leading community-acquired opportunistic mycoses and serious disease (eg, meningitis and cryptococcemia) predominantly occurs in immunocompromised hosts with organ transplantation, hematologic malignancies, or advanced HIV, particularly those not receiving highly active antiretroviral therapy (HAART). Posaconazole has shown good in vitro activity against *C. neoformans* isolates and also inhibited fluconazole-resistant *C. neoformans*. A study of 1,811 global clinical isolates of *C. neoformans* and was much more active than the other triazoles against zygomycetes (Sabatelli et al. 2006). In vitro, posaconazole was highly active against *Candida* spp. The drug is more active than itraconazole and fluconazole against all *Candida* spp. and *C. neoformans*. Posaconazole was the most active azole tested, and was unaffected by fluconazole resistance in *C. albicans* isolates and also had activity against *Candida* spp. that are commonly azole resistant, including *C. glabrata* and *C. krusei*. Posaconazole exhibited fungistatic and fungicidal activity in vitro and in vivo for most *Candida* spp. isolates and inhibited 97% of *Candida* spp. isolates at concentrations of 1 μg/mL or below (Pfaller et al. 2001; Espinel-Ingroff 2003). *C. albicans* is the most susceptible species of *Candida* whereas *C. glabrata* and *C. pelliculosa* are the least susceptible (Pfaller et al. 2001, 2004; Carrillo-Munoz et al. 2005). Nevertheless, posaconazole showed good activity against fluconazole resistant strains especially Candida glabrata, although it was slightly less active than voriconazole (Sabatelli et al. 2006; Pfaller et al. 2008).
showed that of the 1,646 that were tested against posaconazole, 99% had MIC ≤1 μg/mL (Pfaller et al 2005).

Clinical studies
Role of posaconazole as salvage therapy for invasive fungal infections
Salvage therapy of IFI generally refers to the treatment of individuals who are refractory or intolerant to initial therapy, administered for at least 1 week. Besides various case reports several clinical trials of salvage therapy have been undertaken, but most were noncomparator studies or contained a nonrandomized control group. Furthermore response criteria used to document refractory disease differed between these studies. Of note, salvage therapy trials frequently include patients both refractory and intolerant to standard therapy, but patients enrolled because of intolerance have a much better response rate (Maertens et al 2004). When evaluating the results from existing salvage studies one has to take these limitations into consideration.

Invasive aspergillosis
Mortality rates associated with invasive aspergillosis (IA) are still extremely high, particularly in patients undergoing allogeneic HSCT (mortality rate exceeding 50%) and in patients with central nervous system or disseminated infections (Lin et al 2001; Upton et al 2007). A. fumigatus remains the leading cause for these infections, followed by A. flavus, A. niger, and A. terreus, a species that is often resistant to antifungal therapy including amphotericin B. At present, voriconazole is recommended as the primary treatment of IA in most patients (Herbrecht et al 2002; Walsh et al 2008). To date only two clinical externally controlled, open label, multi-center trials have investigated posaconazole as salvage therapy in refractory IA. In the first, 107 patients (initially receiving amphotericin B in most cases) refractory (88%) or intolerant (12%) to conventional therapy were treated with posaconazole (800 mg daily in divided doses) for a median duration of 52 days. Selected subjects of the control group received the best available standard of care for salvage therapy. The overall success rate (defined as complete or partial response) was 42% for posaconazole recipients and 26% for control subjects (Walsh et al 2007). The second study compared 53 patients with hematologic malignancies receiving posaconazole salvage therapy (800 mg daily, median duration 70 days) with 52 contemporary controls treated with high-dose lipid formulation of amphotericin B (HD-LPD/AMB at ≥7.5 mg/kg daily) and with 38 other control patients receiving caspofungin and HD-LPD/AMB. The overall rate of response to posaconazole was 40%, compared with only 8% and 11% for HD-LPD/AMB alone or in combination with caspofungin, respectively. After a follow-up of 12 weeks the overall survival was significantly improved in the posaconazole arm (Raad et al 2008).

In conclusion, these studies indicate that posaconazole is an effective option for salvage therapy for invasive aspergillosis predominantly in patients with hematologic malignancies. Nevertheless, randomized clinical trials to define further the role of posaconazole in IA are warranted, including patients with refractory aspergillosis after first-line therapy with voriconazole.

Oropharyngeal candidiasis in patients with HIV
In patients with HIV and AIDS, oropharyngeal and esophageal candidiasis is the most prevalent opportunistic infection. Fluconazole and itraconazole treatment is usually effective, whereas refractory disease can be observed in 5% of HIV patients, especially in those with advanced HIV infection who have received multiple courses of azole antifungals (Fichtenbaum and Powderly 1998). In the only multi-center, randomized, evaluator-blinded trial of 350 HIV patients with oropharyngeal candidiasis, posaconazole showed noninferiority to fluconazole and a trend to prolonged clinical success after treatment was stopped. Both drugs were administered 14 days (dosed at 200 mg on day one followed by 100 mg daily) and clinical success (evaluated as cure or improvement) was seen in 92% in the posaconazole versus 95% in the fluconazole arm (Vazquez et al 2006). Addressing the role of posaconazole in azole-refractory oropharyngeal and esophageal candidiasis, to date two noncomparative, open-label, multicenter studies demonstrated clinical response rates from 73% to 88%, long-term safety, and good tolerability. Posaconazole was administered 400 mg twice daily, in the first trial (Skiest et al 2007) for a treatment period of 28 days, and up to 12 months in the second trial (Vazquez et al 2007), demonstrating long-term safety, and good tolerability and efficacy.

In summary, posaconazole offers a new, safe, and effective oral treatment option for patients with HIV and azole-refractory mucosal candidiasis.

Zygomycosis
Zygomycosis is a group of frequently lethal mold infections that usually affects diabetic patients and steroid-treated or severely immunocompromised individuals. Most human infections are caused primarily by species of Rhizopus, Mucor, Cunninghamella, Apophysomyces, Absidia, and
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Posaconazole for refractory invasive fungal infections

Rhizomucor. Zygomycetes are highly angioinvasive infections rapidly invading tissue and blood vessels, thereby leading to tissue destruction, thrombosis, infarction, and dissemination. Historically, the agent of choice was conventional amphotericin B at relatively high doses up to 1.5 mg/kg daily, which has largely been replaced by less toxic lipid formulations (Walsh et al 1998; Gleissner et al 2004). Two non-comparative, non-randomized open-label studies of posaconazole for salvage therapy of zygomycosis are available. One report analyzed data from 24 subjects refractory or intolerant to predominantly liposomal amphotericin B as first-line therapy. Median time of posaconazole (800 mg daily in divided doses) administration was 182 days. Complete response, defined as resolution of signs of infection and no relapse within 30 days after cessation of medication, was observed in 46%, and partial response occurred in 33%. Five treatment failures occurred, defined as the presence of zygomycosis at the time of termination of posaconazole treatment or at death. Eighteen patients underwent surgical debridement resulting in a significant better survival (Greenberg et al 2006). In a second study, posaconazole (800 mg daily in divided doses) was evaluated for salvage therapy in 91 cases of probable or proven zygomycosis infection. Pretreatment consisted of amphotericin B in most patients. 81 subjects have failed conventional therapy and 10 showed intolerance or both. Overall, at 12 weeks, 60% of patients had either a complete or partial response to treatment, 17% demonstrated treatment failure, and the remainder had stable disease. Success rates were similar irrespective of site of infection or whether surgical debridement had been performed (van Burik et al 2006).

In refractory zygomycosis, posaconazole constitutes a suitable oral treatment option although to determine its place as first-line therapy a prospective randomized comparison of posaconazole and lipid amphotericin B needs to be performed.

Other invasive fungal infections

Cryptococcal meningitis/fungal CNS infections

Morbidity and mortality associated with fungal infections of the CNS remains unacceptably high despite treatment with a broad variety of antifungal agents. C. neoformans is the most common fungal agent in patients infected with HIV. Existing treatment guidelines for cryptococcal meningitis recommend therapy with amphotericin B with or without flucytosine (Saag et al 2000); however, clinical response rates in HIV-infected individuals are low. In a multi-center, open-label clinical trial, 39 mostly HIV-positive patients with fungal infection of the CNS (29 of them had cryptococcal meningitis) refractory or intolerant to standard therapy with amphotericin B or fluconazole received posaconazole 800 mg daily for a mean duration of 81 days. Successful outcomes were observed in 14 of 29 (48%) subjects with cryptococcal meningitis and in 5 of 10 (50%) patients with CNS infections due to other fungal pathogens (Pitisuttithum et al 2005).

These data suggest that posaconazole has clinical activity against fungal infections of the CNS and provides a valuable alternative in patients failing existing antifungal agents.

Coccidioidomycosis

Coccidioidomycoosis is a systemic fungal infection caused by inhalation of arthroconidia from fungi of the genus Coccidioides. Despite the use of amphotericin B, fluconazole, and itraconazole, disseminated coccidioidomycosis remains difficult to treat, and is characterized by frequent treatment failures and relapses (Galgiani et al 2000). A non-comparative, nonrandomized, multicenter study described 20 patients with chronic pulmonary or nonmeningeal disseminated coccidioidomycosis who received posaconazole (400 mg daily) for up to 6 months. In 85% of treated individuals a satisfactory response, defined as a ≥50% reduction in the Mycoses Study Group score from baseline, could be documented (Catanzaro et al 2007). In a case series of 6 patients with refractory coccidioidomycosis treated with posaconazole (800 mg daily) a successful outcome could be observed in 5 of 6 subjects after 2–34 months of treatment (Anstead et al 2005).

Invasive fusariosis

Fusarium spp. are among the leading fungal pathogens to cause invasive mold infections in patients with underlying hematopoietic malignancy, particularly in those who have undergone HSCT. Conventional amphotericin B-based therapy for invasive fusariosis in patients with hematologic malignancies resulted in a >70% failure rate. Amphotericin B or its lipid formulation represents the treatment of choice, whereas voriconazole is an option for patients refractory or intolerant to first-line therapy. To date one retrospective analysis comprising 21 patients with invasive fusariosis, refractory to or intolerant of standard antifungals (amphotericin B), evaluated posaconazole (800 mg daily in divided doses) as salvage therapy. After a duration of administration for up to 12 months, an overall response rate of 48% could be observed, suggesting that posaconazole is useful as an oral treatment for refractory invasive fusariosis (Raad et al 2006b).
Prophylaxis
Antifungal prophylaxis with posaconazole can be recommended in HSCT recipients with acute graft versus host disease (GVHD) and in patients during induction chemotherapy for acute myelogenous leukemia or myelodysplastic syndrome. These recommendations are based on two randomized clinical studies. One double-blind trial including 600 patients compared posaconazole (600 mg daily in divided doses) with fluconazole (400 mg daily) as prophylaxis against IFI in allogeneic HSCT recipients receiving immunosuppressive therapy for treatment of GVHD. At 4 months, posaconazole was found to be as effective as fluconazole in preventing all IFI (incidence: 5.3% vs 9.0%, p = 0.07) and was superior to fluconazole in preventing proven or probable invasive aspergillosis (2.3% vs 7.0%, p = 0.006). Overall mortality was similar in the two groups, but deaths from IFI were lower in the posaconazole group (Ullmann et al 2007). The second study of 602 subjects compared posaconazole (600 mg daily in divided doses) with fluconazole (400 mg daily) or itraconazole (200 mg twice daily) as prophylaxis for IFI in neutropenic patients receiving chemotherapy for acute myelogenous leukemia or myelodysplastic syndrome (Cornely et al 2007). At 3 months posaconazole was found to be superior to the combined fluconazole/itraconazole group in preventing IFI (2% vs 8%, p < 0.001). Rates of invasive aspergillosis (1% vs 7%, p < 0.001) were also less with posaconazole and a significantly improved survival could be observed (p = 0.04). Posaconazole treatment was relatively well tolerated in both studies.

Safety and tolerability
During phase I, II, and III clinical trials, posaconazole was demonstrated to be safe and well tolerated. In phase I studies most adverse events were mild, transient, and nonspecific, the most common adverse events including gastrointestinal disturbances, headache, dry mouth, somnolence, dizziness, fatigue, and constipation (Courtney et al 2003; Ezzet et al 2005). In a study of 428 patients with refractory invasive fungal infections or febrile neutropenia, 109 of whom continued treatment for ≥6 months, 38% reported adverse reactions (Raad et al 2006a). However, most of these were mild, nausea being the most common (8%), followed by vomiting (6%) and abdominal pain (4%). QT interval prolongation was observed in 1% of patients and elevation of liver enzymes in 2%. Adverse events occurred at similar rates in patients who received posaconazole therapy for <6 months or those treated for <6 months. Long-term therapy did not increase the risk of any individual adverse event, and no unique adverse event was observed with longer exposure to posaconazole (Raad et al 2006a). In two large controlled trials, posaconazole seems to have a comparable safety profile as fluconazole or itraconazole. In a study comprising more than 600 patients comparing posaconazole vs fluconazole or itraconazole prophylaxis in patients with neutropenia, serious adverse events possibly or probably related to treatment were reported by 6% in the posaconazole group and in 2% in the fluconazole or itraconazole group. The most common treatment-related adverse events in both groups were gastrointestinal tract disturbances (Cornely et al 2007; Ullmann et al 2007). Posaconazole therapy is also safe and well tolerated in pediatric and elderly patients and treatment-related discontinuations are uncommon (Torres et al 2005).

Conclusions and place in therapy
Invasive fungal infections cause substantial morbidity and mortality in immunocompromised patients. Since early detection and early effective treatment of invasive fungal infections can be life saving, the most common approach is antifungal prophylaxis and empirical therapy in neutropenic patients with persistent fever. For the future, a more refined approach such as pre-emptive therapy – only initiated upon identification of fungal markers in combination with clinical and radiologic signs – may improve the definition of target patients and expose fewer patients to potentially toxic and/or expensive treatment. The response rate to therapy, in particular for invasive aspergillosis and other invasive mold infections, has been poor. With the advance of a number of techniques facilitating early diagnosis together with the development of new antifungals like the echinocandins or extended spectrum azoles like voriconazole or posaconazole the outcomes for invasive fungal infection in immunocompromised patients could be improved. In this context, caspofungin has emerged as primary therapy of invasive candidiasis (Mora-Duarte et al 2002) and neutropenic fever (Walsh et al 2004) and voriconazole has largely replaced amphotericin B as first line therapy of invasive aspergillosis (Herbrecht et al 2002). Posaconazole is the most broad-spectrum azole antifungal to date, with activity against Candida spp., including fluconazole-resistant isolates, and Aspergillus spp. Posaconazole is likely to be an important new agent in the antifungal armamentarium. The primary role for posaconazole will be the prophylaxis of IFI in severely immunocompromised patients such as HSCT recipients with acute GVHD (Ullmann et al 2007) and neutropenic patients receiving...
intensive induction chemotherapy for acute myelogenous leukaemia and myelodysplastic syndrome (Cornely et al 2007). The European Conference on Infections in Leukemia (ECIL) guidelines therefore included a provisional AI (ie, strongly recommended, based on at least one well-executed, randomized trial) recommendation for these indications (Maertens 2007). However, the prophylactic use of such a broad spectrum agent may lead to the emergence of fungal breakthrough infections as it has been suggested that use of voriconazole as fungal prophylaxis has led to a possible increase of zygomycosis infections in immunocompromised patients. Posaconazole is also approved as first-line treatment of oropharyngeal candidiasis including those refractory to fluconazole or itraconazole in HIV positive patients. There is already clinical evidence for its efficacy as salvage therapy in a number of invasive fungal infections. Posaconazole is the only azole to have activity against the zygomycetes, including *Mucor*, *Rhizopus*, and other species, and has shown clinical effectiveness against refractory infections caused by these fungi. In addition, there are reasonable data showing posaconazole exerts activity against rare, difficult-to-treat fungi such as *Fusarium* and *Scedosporium* spp., and has also activity against *C. neoformans*. Similarly, preliminary evidence from noncomparative salvage trials suggests its efficacy as second-line therapy for invasive *Aspergillus* infection. Nevertheless, many therapeutic areas of uncertainty remain. Amphotericin B and caspofungin remain the only approved agents for the antifungal management of febrile neutropenia, but the broad spectrum of posaconazole makes it an attractive target for further clinical studies. In addition, its role in the treatment of *Candida* spp. remains to be established since no randomized controlled studies in yeast infections are available. Finally, the role of combination therapies of posaconazole with classes of drugs targeting different pathways against filamentous fungi (ie, echinocandins) seems to be a promising task and will provide a focus for future studies.

**Disclosures**

None of the authors has any conflicts of interest to disclose.

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