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Table 1. Fenofibrate Formulations

<table>
<thead>
<tr>
<th>Formulation</th>
<th>Marketed Products</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-micronized</td>
<td>Apo-fenofibrate® 100mg</td>
</tr>
<tr>
<td>Micronized (50% increase in bioavailability compared to non-micronized)</td>
<td>Apo-Feno-Micro® 67mg, 200mg</td>
</tr>
<tr>
<td>Microcoated (75% increase in bioavailability compared to non-micronized)</td>
<td>Lipidil Supra® 100mg, 160mg</td>
</tr>
</tbody>
</table>

The Pharmacy will carry only two formulations of fenofibrate: Apo-fenofibrate® 100mg ($0.36/tablet) and Lipidil Supra® 160mg ($1.33/tablet). Table 2 illustrates the therapeutic interchange policy for fenofibrate orders.

Table 2. Therapeutic Interchange for Fenofibrate

<table>
<thead>
<tr>
<th>Ordered As:</th>
<th>Dispense</th>
</tr>
</thead>
<tbody>
<tr>
<td>Apo-feno-micro® 67mg</td>
<td>Apo-fenofibrate® 100mg</td>
</tr>
<tr>
<td>Fenofibrate 100mg</td>
<td>formulation</td>
</tr>
<tr>
<td>Fenofibrate 200mg</td>
<td></td>
</tr>
<tr>
<td>Lipidil Supra® 100mg</td>
<td></td>
</tr>
<tr>
<td>Fenofibrate 160mg</td>
<td>Lipidil Supra® 160mg</td>
</tr>
<tr>
<td>Lipidil Supra® 160mg</td>
<td>formulation</td>
</tr>
<tr>
<td>Lipidil® 200mg</td>
<td></td>
</tr>
</tbody>
</table>

Deletions

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Any comments, questions or concerns with the content of the newsletter should be directed to the editors. Write to CSU Pharmaceutical Sciences, Vancouver General Hospital, 855 W12th Ave, Vancouver BC V5Z 1M9, send a FAX to 604-875-5267 or email kshalans@vanhosp.bc.ca
Find us on the Web at www.vhpharmsci.com
Deletions

1. Pindolol tablets (Visken®)
   - Alternative: Acebutolol

2. Gemfibrozil capsules (Lopid®)
   - Alternative: Fenofibrate

Updated Policies

1. Serum Drug Level Monitoring

Clinical pharmacists have been authorized to order serum concentrations of all measurable drugs including (but not limited to) aminoglycosides (with serum creatinine), carbamazepine, cyclosporine, digoxin, lithium, phenobarbital, phenytoin (with serum albumin), tacrolimus, theophylline, valproic acid, and vancomycin (with serum creatinine).

2. Revised Drug Administration Policies

The on-line and hard copy versions of the Parenteral Drug Therapy Manual (PDTM) have been updated with these new policies:

- Skeletal muscle relaxants (cisatracurium, pancuronium, rocuronium, succinylcholine) are restricted to critical care areas.

- The following platelet inhibitors: abciximab, eptifibatide and tirofiban may be administered in any critical care area.

New Drugs/Drug Products

1. Tiotropium inhaler (Spiriva®)

   Jennifer Haymond B.Sc. (Pharm), Karen Shalansky, Pharm.D.

   Tiotropium is a long-acting anticholinergic agent indicated for long-term maintenance treatment of bronchospasm associated with chronic obstructive pulmonary disease (COPD).  

Pharmacology

Tiotropium is a quaternary ammonium muscarinic receptor antagonist demonstrating specificity for the M3-receptor subtype. The M3-muscarinic receptor is largely responsible for visceral smooth muscle contraction in response to acetylcholine. By antagonizing the effects of acetylcholine on bronchial smooth muscle, tiotropium reduces bronchospasm. Other proposed benefits of anticholinergic bronchodilators include regulating mucus secretion and inhibiting viral influence on cholinergic tone.

Tiotropium demonstrates a long duration of action of at least 24 hours that can be attributed to its high affinity for and slow dissociation from the M3-muscarinic receptor.

Comparable Formulary Bronchodilator Agents

Ipratropium, a short-acting anticholinergic bronchodilator, and salmeterol and salbutamol, long-acting and short-acting beta2 (β2) agonist bronchodilators, respectively, are currently on formulary (Table 3).

<table>
<thead>
<tr>
<th>Drug Name</th>
<th>Mechanism of Action</th>
<th>Time to Onset (minutes)</th>
<th>Duration of Effect (hours)</th>
<th>Typical Dose</th>
<th>VH Acquisition Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tiotropium (Spiriva®)</td>
<td>Anticholinergic; antagonizes M3-muscarinic receptors</td>
<td>30</td>
<td>24</td>
<td>18 mcg once daily</td>
<td>$10.50/5 x 18 mcg capsules Inhaler device free</td>
</tr>
<tr>
<td>Ipratropium (Atrovent®)</td>
<td>Anticholinergic; non-selective blocker of muscarinic receptors</td>
<td>5-15</td>
<td>4-6</td>
<td>40mcg QID</td>
<td>$12.79/20mcg inhaler (140-dose inhaler) (nebulizer available)</td>
</tr>
<tr>
<td>Salmeterol (Serevent®)</td>
<td>Beta2 agonist; ↑ cyclic adenosine monophosphate (cAMP)</td>
<td>10-20</td>
<td>12-18</td>
<td>50mcg BID</td>
<td>$24.90/25mcg inhaler (120-dose inhaler)</td>
</tr>
<tr>
<td>Salbutamol (Ventolin®)</td>
<td>As above</td>
<td>2-5</td>
<td>4-6</td>
<td>200mcg q4-6h</td>
<td>$2.65/100mcg inhaler (200-dose inhaler) (nebulizer available)</td>
</tr>
</tbody>
</table>

Table 3. Comparison of Formulary Bronchodilators for COPD
Compared to placebo, tiotropium provides superior bronchodilation and quality of life scores, and fewer exacerbations and hospitalizations in patients with COPD. There are two identical randomized, double-blind, parallel group studies directly comparing tiotropium to ipratropium. Results of the two studies were combined after one year. Patients received either tiotropium 18 mcg daily (n=356) or ipratropium 40mcg qid (n=179). Tiotropium demonstrated significantly greater improvement in trough FEV₁ and FVC throughout the study. While both groups used lower rescue salbutamol compared to baseline, the reduction was greatest in the tiotropium group (p<0.05 for 40 of the 52 weeks). Tiotropium significantly reduced the proportion of patients with one or greater COPD exacerbations (35% vs. 46%, p=0.014) although the proportion of hospitalized patients was similar (7.3% vs. 11.7%, p=0.11).

Trials comparing tiotropium to salmeterol have been unable to demonstrate significant differences in reducing exacerbation rates or use of rescue salbutamol in patients with moderate to severe COPD. However, tiotropium demonstrated significantly improved FEV₁ over salmeterol at 24 weeks (p<0.01). This difference may be attributed to a possible tachyphylactic effect with prolonged salmeterol use, resulting in a decrease in salmeterol response.

Potential Risks
The most commonly reported adverse event for tiotropium in clinical trials is dry mouth. In the 1-year comparative trial of tiotropium and ipratropium, dry mouth was significantly higher in the tiotropium group (12.1% vs. 6.1%, p=0.03). Dry mouth is usually mild and has led to a 0.3% discontinuation rate.

Potential Benefits/Role in COPD Therapy
Tiotropium has demonstrated improvement in reducing exacerbation rates compared with placebo (RR 0.74; 95% CI 0.62-0.89) or ipratropium (RR 0.78; 95% CI 0.63-0.95). Additionally, tiotropium has demonstrated consistent, significant improvements in reducing exacerbation and hospital admission rates over placebo whereas salmeterol has not. Once daily administration is also advantageous for compliance purposes and for elderly patients who have difficulty manipulating a metered dose inhaler.

The 2003 guidelines from the Canadian Thoracic Society recommend a long-acting bronchodilator as the second step in therapy if COPD symptoms persist despite reasonable short-acting bronchodilator use (level of evidence: 1A). Recommended long-acting bronchodilators include tiotropium or a long-acting β₂ agonist. Combination tiotropium and a long-acting β₂ agonist are recommended for patients with persistent moderate to severe symptoms (level of evidence: 3A).

Dose and Cost
The dose of tiotropium is inhalation of one capsule (18mcg) once daily using the supplied handihaler. Tiotropium capsules come in packages of 10 that can be split into two, allowing a 5-day course of tiotropium to be dispensed ($10.50/5 days). The inhaler itself is provided free of charge to the hospital.

Conclusion
Tiotropium demonstrates superior bronchodilation compared to placebo and ipratropium. Studies have yet to demonstrate significant superiority of tiotropium over salmeterol. Tiotropium is currently recommended in the Canadian COPD Guidelines as an option for patients who are not optimally controlled with short-acting bronchodilators. Ipratropium is still required on formulary due to its availability as a nebulizer for use in acute asthma/COPD exacerbations and as well for initial therapy of mild COPD.

References
Bisoprolol (Monocor®)
Karmin Ip B.Sc.(Pharm), Susan Buchkowsky B.Sc.(Pharm), Rubina Sunderji, Pharm.D.

Bisoprolol is a $\beta_1$-adrenergic receptor blocker (cardioselective beta-blocker) indicated for the management of hypertension and heart failure (HF). It does not have intrinsic sympathomimetic activity or membrane stabilizing activity

Clinical Trials
Two randomized placebo-controlled trials evaluating bisoprolol in patients with HF have been published: CIBIS I and CIBIS II (Table 4). In the CIBIS trial, patients continued to receive standard therapy with an angiotensin converting enzyme inhibitor (ACEI) (90%) and a diuretic (100%) and were followed for a mean of 1.9 years. The observed difference in mortality in favour of bisoprolol did not reach statistical significance (16.6% vs. 20.9%). The inability to detect a survival difference was attributed to a lower than expected mortality rate used in the power calculation and possible suboptimal dosing of bisoprolol (target 5mg/day).

The CIBIS II trial was designed using a revised estimate of placebo mortality rate from the CIBIS trial and a higher bisoprolol dose (target 10mg/day). In this larger trial, patients were randomized to placebo or bisoprolol on background therapy with diuretics (98.5%) and ACEI (96%) and were followed for a mean of 1.3 years. The trial was stopped early because the all-cause mortality was significantly less in the bisoprolol group compared to placebo (11.8% vs. 17.3%, p < 0.0001). The bisoprolol group also had significantly fewer cardiovascular deaths (9% vs 12%, p=0.004) and hospital admissions (33% vs. 39%, p=0.0006). A subgroup analysis of the CIBIS II trial examining the dose-response of bisoprolol showed a similar reduction in mortality with low, moderate and high doses of the drug.

In both CIBIS trials, bisoprolol was well tolerated with no significant difference in withdrawal rates due to adverse effects compared to placebo.

Comparison with other Beta-Blockers for HF
Only two other beta-blockers, metoprolol and carvedilol have shown survival benefit in HF in randomized, placebo-controlled trials. Metoprolol is cardioselective whereas carvedilol is a non-selective agent with $\alpha_1$, $\beta_1$, and $\beta_2$ blocking properties (Table 5). The alpha blockade with carvedilol has resulted in a higher incidence of side effects such as dizziness (32.4% carvedilol vs. 19.2% placebo) in clinical trials.

Similar to bisoprolol, metoprolol and carvedilol have also shown to reduce mortality compared to placebo in HF (Table 4). In a head-to-head trial, carvedilol was significantly superior to regular release metoprolol (COMET). Of note, the formulation of metoprolol that improved survival in the MERIT-HF trial is a controlled release preparation (metoprolol XL) that is not available in Canada. It is speculated that the controlled release preparations lead to a more pronounced and even beta-blockade over 24 hours in comparison to regular release preparations. This degree of blockade may be important in protecting the heart from abrupt increases in sympathetic nervous activity which may potentiate ventricular fibrillation and sudden death, and in long-term myocardial performance of the failing heart.

Dose and Cost
The initial dose of bisoprolol is 1.25mg daily gradually increased by 1.25mg every 1-4 weeks to a maximum target dose of 10mg daily for HF.

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**Table 4. Studies of Beta-Blockers in Heart Failure**

<table>
<thead>
<tr>
<th>Study (Drug)</th>
<th>Design (N)</th>
<th>Population</th>
<th>Target Dose</th>
<th>Mortality</th>
</tr>
</thead>
<tbody>
<tr>
<td>CIBIS I (Bisoprolol)</td>
<td>R, DB (641)</td>
<td>EF ≤ 40% NYHA III-IV</td>
<td>5mg/day</td>
<td>B:16% P: 20.9% (NS)</td>
</tr>
<tr>
<td>CIBIS II (Bisoprolol)</td>
<td>R, DB (2547)</td>
<td>EF ≤ 35% NYHA III-IV</td>
<td>10mg/day</td>
<td>B:11.8% P: 17.3% (p&lt;0.0001)</td>
</tr>
<tr>
<td>MERIT-HF (Metoprolol XL)</td>
<td>R, DB (3991)</td>
<td>EF ≤ 40% NYHA II-IV</td>
<td>200mg/day</td>
<td>M:7.3% P: 10.8% (p=0.0062)</td>
</tr>
<tr>
<td>US Carvedilol</td>
<td>R, DB (1094)</td>
<td>EF ≤ 35% NYHA III-IV</td>
<td>50mg BID</td>
<td>C:3.2% P:7.8% (p&lt;0.001)</td>
</tr>
<tr>
<td>COPERNICUS (Carvedilol)</td>
<td>R, DB (2289)</td>
<td>EF ≤ 25% NYHA IV</td>
<td>25mg BID</td>
<td>C:11.2% P:16.8% (p=0.0014)</td>
</tr>
<tr>
<td>COMET (Metoprolol vs Carvedilol)</td>
<td>R, DB (3029)</td>
<td>EF &lt; 35% NYHA II-IV</td>
<td>M: 50mg BID C: 25mg BID</td>
<td>M:40% C:34% (p=0.0017)</td>
</tr>
</tbody>
</table>

R=Randomized, DB=Double Blind; EF=Ejection Fraction; NYHA=New York Heart Association; B=Bisoprolol; P=Placebo; M=Metoprolol; C=Carvedilol; NS=not significant.
Table 2 illustrates equivalent doses and costs of various beta-blockers including bisoprolol. Both bisoprolol and metoprolol are Pharmacare benefits. Carvedilol 25-50mg bid is the most costly of the three agents and requires a completed special authority form for Pharmacare benefit.

Conclusion
Bisoprolol has been added to formulary for the following reasons:
- Unlike regular release metoprolol, bisoprolol has proven to decrease mortality from HF in a large randomized trial.
- Bisoprolol is preferred over carvedilol due to its lower cost and does not require special authority for Pharmacare benefit.
- Bisoprolol's pharmacokinetic profile allows for once daily dosing and improved patient compliance.

References

Residency Accreditation Status
The Vancouver Hospital Pharmacy Residency Program was recently awarded full accreditation by the Canadian Hospital Residency Board. Congratulations to Dr. Kerry Wilbur and Dr. Peter Loewen who are the Residency Program co-coordinators and all the other contributors to the residency program.