Cost-effectiveness of treprostinil versus epoprostenol in patients with pulmonary arterial hypertension: A Canadian analysis

Thomas R Einarson PhD1,2, John T Granton MD FRCS3, Colin Vicente BSc2, John H Walker PhD2,4, Greg Engel BSc5, Michael Iskedjian BPharm MSc2

BACKGROUND: Pulmonary arterial hypertension (PAH) is associated with substantial morbidity and mortality, exerting a tremendous health and economic impact on patients. In the present study, an economic evaluation of patients with PAH treated with either treprostinil or epoprostenol was performed.

METHODS: A cost-minimization analysis (a cost-effectiveness subtype) was performed under the assumption that treprostinil and epoprostenol were clinically equivalent. Two cohorts of 60 patients, treated with treprostinil or epoprostenol, were evaluated over three years by using a dynamic spreadsheet model. The evaluation included both the provincial ministries of health and societal perspectives. Resource valuation data for drugs, medical supplies, consultations, and surgical and diagnostic procedures were obtained from standard lists. Costs of hospitalizations and adverse events were derived from published sources. Additional outpatient costs were considered equivalent and, therefore, were excluded from the analysis. Costs are presented in 2003 Canadian dollars discounted at 3%. Sensitivity analyses were performed testing all uncertainties in the model.

RESULTS: In the base-case analysis (over three years), treatment with treprostinil resulted in an expected savings of $2,610,642 and $2,781,438 from the ministries of health and societal perspectives, respectively. On a per-patient level, treatment with treprostinil resulted in an average annual savings of $14,504 and $15,452, respectively. The greatest savings with treprostinil came from reduced hospitalizations. Multivariate sensitivity analyses estimated cost savings in greater than 99% of scenarios.

CONCLUSIONS: By initiating and continuing treprostinil treatment over a three-year period, the economic burden associated with PAH may be reduced compared with epoprostenol treatment.

Key Words: Cost-effectiveness; Epoprostenol; Health economics; Pulmonary arterial hypertension; Treprostinil

Pulmonary arterial hypertension (PAH) is a condition of abnormally high pressure within the pulmonary arterial circulation, characterized by an increase in pulmonary vascular resistance and a mean pulmonary artery pressure of 25 mmHg or greater (1,2). No investigations to date have provided definitive data on the incidence, prevalence and mortality of PAH.

However, idiopathic PAH (IPAH), a subtype of PAH, is estimated to have an annual incidence of one to two cases per one million people in both the United States and Europe (3). Based on that rate, an estimated 300 to 600 new cases of IPAH would occur per year in the United States, or 30 to 60 new cases in Canada. Left untreated, IPAH leads to premature death.

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one study from the United States (4), the mean survival time was 2.8 years, and the five-year survival rate was 34%. Therefore, IPAH is a disease with serious implications.

The treatment of IPAH is aimed at relieving symptoms, improving physical activity and increasing survival (1). Prostacyclins constituted the first therapeutic class of drugs approved by Health Canada for the treatment of severe New York Heart Association (NYHA) functional class III and class IV pulmonary hypertension. Flolan (epoprostenol; GlaxoSmithKline, Canada), the first approved prostacyclin, is indicated for the long-term intravenous treatment of IPAH and secondary pulmonary hypertension due to the scleroderma spectrum of diseases in NYHA functional class III and class IV patients who are not adequately responding to conventional therapy (5,6). Unfortunately, because epoprostenol must be delivered as a continuous intravenous infusion, it may be associated with various severe adverse events such as septicemia and the threat of acute hemodynamic instability from interrupted line flow due to the loss of line integrity (7).

In October 2002, United Therapeutics (USA) received approval from Health Canada to market Remodulin (treprostinil sodium), a new prostacyclin for the long-term, subcutaneous treatment of PAH in NYHA class III and class IV patients who are not adequately responding to conventional therapy (8). Treprostinil is an improvement over previous prostacyclins; it has an increased half-life and has been shown to be an effective treatment when administered via a continuous subcutaneous infusion (9,10). The subcutaneous route of delivery results in fewer serious adverse events than the intravenous route. These serious adverse events include local and systemic infections, thrombosis and paradoxical emboli (11). Furthermore, the subcutaneous route of delivery may improve patient survival compared with conventional therapy (12). Results from a recent clinical trial (13) suggest that three-year patient survival with subcutaneously administered treprostinil is no worse than patient survival with intravenously administered epoprostenol.

The differences between the delivery routes of the two drugs are expected to have an impact on the rate of medical resource utilization required for the clinical management of PAH. In the United States, Highland et al (14) performed a decision analysis examining the cost-effectiveness of bosentan, epoprostenol and treprostinil in treating PAH. The study reported that over a one-year treatment horizon, treatment with bosentan would provide a cost savings compared with both treprostinil and epoprostenol. In addition, based on various assumptions, they reported that treatment with epoprostenol may provide a cost savings compared with treprostinil.

To date, no Canadian study has assessed this difference in terms of an economic evaluation of the two drugs. The present study is the first to examine this issue, and the first to report a pharmacoeconomic evaluation of subcutaneous treprostinil compared with intravenous epoprostenol in the treatment of patients with NYHA class III or IV PAH from the viewpoint of the Canadian health care system.

METHODS

The primary audience for the present evaluation includes the various Canadian provincial drug benefit plans as payers, physicians as prescribers, and patients as users. The primary analytical perspective was that of the various Canadian provincial ministries of health (MoH). Secondary perspectives included the societal perspective to provide results to the decision makers of the various benefit plans.

A three-year time horizon was chosen for the analysis based on a median survival time of 2.8 years in patients untreated for IPAH (4). This time horizon allowed the capture of all relevant outcomes in the decision model. The model incorporated the dose-titration phase in addition to maintenance therapy.

In accordance with the Canadian and Ontario guidelines for economic evaluations of pharmaceuticals, the pharmacotherapy being evaluated was compared with the existing practice (ie, the most prevalent clinical practice, and either the lowest cost comparator or the ‘do-nothing’ approach) (15,16). In the case of treprostinil, the most appropriate comparator was epoprostenol. Due to the severe nature of the disease, the do-nothing approach was not considered appropriate.

Model design

The analyses were conducted by using a decision analytical spreadsheet model based in Microsoft Excel (Microsoft Corporation, USA). The economic model used was a cost-minimization analysis. This model was adopted as a result of preliminary data analysis, expert clinical opinion, and results from noncomparative studies indicating that both treprostinil and the comparator, epoprostenol, provide an effect significantly greater than placebo. In addition, based on the results of a recent three-year clinical trial (13), the two drugs provide equal survival in patients with severe PAH.

An incidence-based model was used to follow both treprostinil and epoprostenol treatment in two cohorts of 60 patients with PAH in NYHA class III and class IV. Each patient underwent a dose-titration phase in addition to maintenance therapy, as determined by specialists experienced in managing such cases. The spreadsheet model was designed to represent the logical sequence of clinical practice as described in the literature. All points were verified by expert clinical opinion.

Resource utilization

Parameter values for the analyses were primarily determined from the published literature. Resource utilization data were derived from clinical trials and published treatment guidelines. Expert clinical opinion was used to confirm the resource utilization data extracted and to provide guidance in areas of uncertainty requiring further sensitivity analyses.

Drug costs were determined from standard lists wherever available (eg, the Ontario Drug Benefits Formulary or the Régie de l'Assurance-médicaments du Québec) (17-19). The unit cost of treprostinil was provided by the manufacturer as a provisional cost within a plausible range; this range was applied to sensitivity analyses. Treprostinil is administered through a continuous subcutaneous infusion and, for the present analysis, was dosed based on a 0.82:1 ratio versus epoprostenol (20). The average dose used in the base-case analysis was 22.2 ng/kg/min (daily dose of 2.24 mg). The unit cost of treprostinil used was $45.00/mg. Because treprostinil is initiated as an outpatient therapy in the majority of cases, the dose is titrated over a period of two to 12 weeks to reach a therapeutic dose level based on the patient’s response and ability to tolerate titration. The inpatient daily dose cost used was $100.08 because the delivery system was a multidose vial without any drug wastage.

Epoprostenol is administered by intravenous infusion and usually requires hospitalization for a period of five to 10 days to titrate the drug to a maintenance level and to ensure enough time for the patient to receive proper training on how to reconstitute the drug and how to appropriately care for the drug delivery system. Treatment is
normally initiated at 2 ng/kg/min to 4 ng/kg/min with a target dose of 10 ng/kg/min to 15 ng/kg/min in two to four weeks (21). The mean dose for adults is approximately 20 ng/kg/min to 40 ng/kg/min (22). In the present analysis, the mean weight of a patient was estimated to be 70 kg and the mean dose was 27.2 ng/kg/min (daily dose of 2.72 mg) (20). The inpatient daily dose cost of epoprostenol was $83.49 (Table 1).

Daily medication costs were determined based on the number of units required for each daily dose. In-hospital medication costs did not include any additional dispensing fees. However, outpatient medication costs included a $6.54 dispensing fee for each prescription and a 10% markup on the cost of the drug, less a $2.00 copayment. These drug costing approaches were used for both treprostinil and epoprostenol.

Medical consultation fees were derived from the Ontario Schedule of Benefits for Physician Services (18). All patients incurred the cost of an initial consultation with their family practitioner and the cost of referral to a specialist, regardless of the treatment. Experts were consulted to establish the type and frequency of visits that patients would require for each drug. For both treatments, a team of specialists, including a cardiologist, rheumatologist and a respiratory disease specialist, initially examined each patient. Although patients should be seen by the entire consulting team on a monthly basis, it was assumed that some patients would be unable to make monthly visits. As a result, it was estimated that 50% of patients would adhere to the monthly visits and 50% would make bimonthly visits. Because clinical practice may differ dramatically among practitioners, and some may only follow-up on a patient every three months or longer, it was preferable to be conservative and err toward a higher frequency of visits.

In addition to visits for specialist consultations, medical visits included the nurse’s time to train the patients. Before discharging patients whose drug dose was being titrated, both the patient and a spouse or close relative received training from a specially trained nurse on proper catheter care, sterile techniques and drug preparation and administration.

Costs of surgical and diagnostic procedures were derived from the Schedule of Benefits for Physician Services and other standard price reference lists such as the Ontario Schedule of Benefits for Laboratory Services (5,17). Epoprostenol requires the surgical placement of an intravenous catheter (treprostinil does not require such a procedure); thus, in the model, all patients treated with epoprostenol incurred the cost of having this procedure performed. Total costs of other diagnostic procedures were based on the frequency of performing each assessment and the proportion of patients who required the procedure, with data being derived from the literature (2,23). Utilization rates were verified by expert clinical opinion.

Hospitalization rates and utilization were determined from the literature (2,24). Each listed reference presented data from several sources. Costs varied slightly according to the source; however, the valuations were similar among all reported resources.

Epoprostenol use, because of its route of infusion, puts patients at risk for sepsis and line infections (5). McLaughlin et al (7) reported a 14% rate of sepsis for epoprostenol treatment. For this reason, the present study also sought to determine the cost of treating sepsis. The costs of treating other adverse events associated with treprostinil or epoprostenol were considered comparable or minor; hence, they were not considered in the present analysis. For example, the cost to manage injection site pain in treprostinil patients was not included. However, injection site pain is presumably managed by outpatient treatment, and it was not expected to substantially affect the results of the analysis.

### TABLE 1

| Medication           | Utilization* | Valuation ($) |
|----------------------|--------------|---------------|
| **Trexolint** 22 ng/kg/min | 50 mL/inl | 45.00/mg |
| Epoprostenol§ 27 ng/kg/min | 50 mL/inl | 23.33/mg |
| Drug diluent§ 50 mL/vial | 50 mL/inl | 10.00/vial |

### Infusion pump and supplies

| Medication       | Valuation ($) |
|------------------|---------------|
| **Trexolint** 100% | 21.00/day |
| Epoprostenol§ 100% | 44.00/day |

### Physician visits§

| Medication       | Valuation ($) |
|------------------|---------------|
| General physician | Yrly 54.75   |
| Cardiologist, rheumatologist, and respiratory specialist | Yrly 125.00 |
| Consultation Subsequent visits 50% – monthly | 24.65 |
| Subsequent visits 50% – bimonthly | 24.65 |
| Training nurse** | Treprost. – 1 session 30.00/day |
| Epoprostenol 5 sessions | 20.00/day |

### Diagnostic procedures††

| Medication      | Valuation ($) |
|-----------------|---------------|
| Echocardiogram  Every 6 months | 12.55 |
| Right heart catheterization Every 6 months | 151.10 |
| Pulmonary angiogram Every 6 months | 107.50 |
| Exercise stress test Every 6 months | 107.80 |
| Chest radiography Yrly 39.50 |
| Thorax CT Yrly 65.60 |
| Oxygen consumption studies | 16.80 |
| Blood cultures – CBC Yrly 7.75 |
| Blood-gas analysis Every 6 months | 11.45 |
| Lung compliance Every 6 months | 99.95 |
| Simple spirometry Every 6 months | 4.25 |
| Oxygen saturation Every 6 months | 4.25 |

### Surgical procedures‡‡

| Medication       | Valuation ($) |
|------------------|---------------|
| Hickman or Broviac CVC§§ 0% – treprostinil | 132.85 |
| Removal of CVC 0% – epoprostenol | 38.70 |

### RESOURCE VALUATION

Resource utilization and the unit costs of medications, delivery systems, consultations and laboratory and diagnostic procedures are presented in Table 1. For epoprostenol, the unit cost per milligram was determined to be $23.33 (19). However, the daily medication cost of epoprostenol also included the cost of the diluent used to prepare the medication. The cost per vial of diluent required for the constitution of Flolan was $10.00 (19). Coverage rates for both the drug delivery system and required infusion supplies were determined to be $21 per day and $44 per day for treprostinil and epoprostenol, respectively. The utilization rates for procedures and diagnostics derived from expert opinion were lower than the frequency suggested in the guidelines. However, variations in utilization rates were equal in both arms. Thus, any bias introduced would have been the same for both treatment arms.

**Data from references 2 and 23 (all points were verified by expert clinical opinion); †The source of valuation was the manufacturer; ‡Data from reference 20; §Data from reference 19; ¶Data from reference 5; **Data from reference 26; ††Data from references 5 and 17; ‡‡Manufactured by Bard Access Systems, USA. CBC Complete blood count; CT Computed tomography; CVC Central venous catheter
Multivariate sensitivity analyses were performed by using Monte Carlo simulations to determine the robustness of our model to probability distributions for each uncertain parameter. The simulations were conducted by using 10,000 iterations for each analysis. The results of each analysis are presented as the mean value of the results generated from the 10,000 iterations.

### Sensitivity analyses

One-way sensitivity analyses were performed to determine the impact of varying specific parameters. The sensitivity of the model to the dose adjustment factor was tested by varying the dose ratio while maintaining all other variables constant. The cost savings of treprostinil were expected to be reduced as the dose ratio approached 1.09:1. Results of the sensitivity analyses are shown in Table 6. A threshold analysis was also performed to determine the dose ratios at which point there would be no more savings with treprostinil versus epoprostenol treatment. The break even point was determined to be a single 15 min session on training patients was determined to be a single 15 min session for treprostinil and five daily 15 min sessions for epoprostenol.

To determine the total cost of hospitalization, the average length of stay (ALOS) in the hospital was multiplied by the average daily hospital cost. The ALOS reported in the literature for epoprostenol treatment was 15 days per rehospitalization for NYHA class III and class IV patients, and was confirmed by expert clinical opinion (25). A summary of the utilization of hospital resources is presented in Table 2. To standardize the published hospital costs (26,27) from previous years into 2003 Canadian dollars, published dollar values were adjusted by using the health portion of the consumer price index (28). Hospital costs used in the model are presented in Table 3.

### Sensitivity analyses

Various assumptions were used to facilitate the analysis. Each assumption was determined by either a literature review or from expert clinical opinion. The uncertainty inherent in each assumption was tested by sensitivity analyses, except for the assumption that the frequency of patient consultations was equally distributed over both comparators. Because this assumption did not introduce any bias, it was not expected to influence the results.

The sensitivity analyses are presented in Table 6. A threshold analysis was also performed to determine the dose ratios at which point there would be no more savings with treprostinil versus epoprostenol treatment. The break even point was determined at a dose ratio of 1.48:1 at one year and 1.20:1 at three years. One-way sensitivity analyses were performed from both the MoH and societal perspectives. The probability of cost savings with treprostinil. The break even point was determined at a dose ratio of 1.48:1 at one year and 1.20:1 at three years. One-way sensitivity analyses were performed from both the MoH and societal perspectives. The probability of cost savings with treprostinil versus epoprostenol treatment was 100% in the first year and greater than 99% over three years from both the MoH and societal perspectives. Results of the multivariate analyses are presented in Table 7. The results indicate robustness against alterations in all important parameters across all plausible ranges.
DISCUSSION

The present economic evaluation was performed to provide the decision makers of various drug plans with a pharmacoeconomic profile of treprostinil in the treatment of patients with severe PAH (NYHA class III or IV). In the base-case analysis, the first three years of treating the cohort with treprostinil resulted in a cost savings of $2,610,642 and $2,781,438 from the MoH and societal perspectives, respectively. On an individual basis, the per-patient incremental cost savings were $43,511 and $43,357 from the MoH and societal perspectives, respectively. Thus, treprostinil appears to be cost saving compared with epoprostenol.

To examine the impact of varying the values of key model parameters, a number of one-way sensitivity analyses were performed. The model results were sensitive to scenarios including varying the dose adjustment ratio. In the majority of cases, most of the model results were congruent in magnitude with the cost savings found from the base-case analysis. The Monte Carlo simulations predicted results similar to what was determined through one-way sensitivity analyses. Cost savings after the third year of treatment were favourable for treprostinil, with cost savings in more than 99% of simulations from both the MoH and societal perspectives.

No other Canadian economic evaluation of treprostinil has been published. An American study by Highland et al (14) reported results that were inconsistent with our findings.

TABLE 4
Parameter values tested in the multivariate sensitivity analyses

| Parameter | Base-case value (sensitivity) | Distribution | Reason for range |
|-----------|------------------------------|--------------|------------------|
| Discount rate | 3% (0–5) | Normal | As per CCOHTA guidelines* |
| Dose adjustment factor | 0.82:1 (0.82:1–0.91:1) | Triangular | As per rates reported in the literature† |
| Rate for treprostinil delivery system | $21/day (20–25) | Triangular | Range provided by expert clinical opinion |
| Duration of hospitalization to titrate medication | 0 days (0–2) | Exponential | Expert opinion |
| Epoprostenol | 10.5 days (7–14) | Normal | Expert opinion |
| Duration of training | 1 day (1–2) | Poisson | Expert opinion |
| Epoprostenol | 17.5 days (14–21) | Poisson | Expert opinion |
| Rate of sepsis | 0.14/person-year (0.08–0.32) | Log normal | High end reported in product monograph‡ |
| Cost to treat sepsis | $16,228/episode (8114–32,456) | Triangular | Estimated range of −50% to +200% based on skewness of cost data |
| Rate and length of stay for rehospitalizations | 10.2 days/year (3.4–37.4) | Poisson | As per standard deviation reported in prospective economic evaluation§ |
| Treprostinil | 15 days/year (5–55) | Poisson | Expert opinion |
| Epoprostenol | 8 h/day hospitalized (8–24) | Uniform | Range based on average work day of 8 h to entire 24 h spent in hospital per hospital day |
| Duration of lost time for hospitalizations | $17.18/h (17.18–34.36) | Triangular | Range based on average hourly wage to ‘double-time’ |

*Data from reference 16; †Data from references 20 and 30; ‡Data from reference 5; §Data from reference 25. CCOHTA Canadian Coordinating Office for Health Technology Assessment

TABLE 5
Itemized costs of each therapy

| Item | Cohort | Drug and cost | 202,985 | 2,234,806 | 5,666,759 | 162,923 | 1,847,786 | 4,693,427 |
|------|--------|---------------|--------|-----------|-----------|--------|-----------|-----------|
| Item | Equipment | 38,351 | 422,236 | 1,070,657 | 80,355 | 884,685 | 2,243,281 |
| Item | Medical visits | 28,463 | 68,844 | 98,694 | 69,341 | 112,590 | 142,440 |
| Item | Procedures | 37,455 | 64,367 | 160,372 | 45,426 | 74,112 | 184,620 |
| Item | Hospitalization | 120,211 | 770,092 | 1,870,521 | 1,375,377 | 2,289,579 | 3,896,749 |
| Item | AE – sepsis | 0 | 0 | 0 | 11,360 | 125,066 | 317,127 |
| Item | Total MoH cost | 427,465 | 3,560,345 | 8,867,003 | 1,744,782 | 5,333,819 | 11,477,645 |
| Item | Lost time | 11,181 | 81,708 | 201,039 | 96,895 | 197,553 | 371,835 |
| Item | Total SOC cost | 438,646 | 3,642,053 | 9,068,042 | 1,841,677 | 5,531,372 | 11,849,480 |

AE Adverse events; MoH Ministry of health perspective; SOC Societal perspective

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However, various limitations were present in that study, such as the assumption of equal hospital resource utilization between the two therapies and the use of average doses that are out of line with current dose studies (20). In addition, the decision analysis performed did not include the cost to treat sepsis, as indicated in both the product monograph and recent epoprostenol studies (5,7).

Although no comparable Canadian cost-effectiveness study was available, an unpublished cost analysis of epoprostenol in 27 Canadian patients with PAH did report a trend in epoprostenol resource valuation that was similar to that in our study (24). The average cost per day of epoprostenol treatment, including the direct and indirect costs reported in that study, was $211 (2001 Canadian dollars). In the present analysis, we concluded that the average cost per day of epoprostenol was $135 from the MoH perspective and $175 from the societal perspective. As a result, our economic evaluation produced conservative expected costs of epoprostenol compared with the previous prospective cost analysis; however, the relative impact of the frequency of follow-up visits would not change the overall direction of the results. Additionally, the base-case analysis did not include the cost of pain management for each treatment. The actual cost of pain management could not be captured accurately and, therefore, it did not seem sensible to include the assumption in our base-case analysis, especially from the MoH perspective.

**CONCLUSIONS**

Although treprostinil and epoprostenol have shown comparable clinical efficacy, they are administered by different delivery systems. The treprostinil delivery system has been associated with an improved safety profile compared with epoprostenol. As a result, the economic profile of treprostinil was expected to be more resource efficient than that of epoprostenol.

As our results suggest, based on a cost-minimization analysis, treprostinil is a more resource efficient treatment, providing a cost savings over epoprostenol of $43,511 per patient over a three-year period. The greatest cost savings were attributed to a decrease in the ALOS during dose titration and fewer

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**TABLE 6**

| Scenario | MoH ($) | Societal ($) |
|----------|---------|--------------|
| Base-case savings | 2,610,642 | 2,781,438 |
| Discount rate, 0% to 5% | 2,726,857 to 2,538,346 | 2,905,264 to 2,704,407 |
| Dose adjustment ratio, 0.82:1 to 1:09:1 | 2,610,642 to 747,191 | 2,781,438 to 917,987 |
| Sepsis rate, 0.08 to 0.32 events per person-year | 2,311,259 to 3,508,788 | 2,464,789 to 3,731,383 |
| Average cost per episode of sepsis, $8114 to $32,456 | 2,452,078 to 2,927,769 | 2,622,874 to 3,098,565 |
| ALOS during dose titration | | |
| Treprostinil, 0 to 2 days | 2,610,642 to 2,337,120 | 2,781,438 to 2,491,904 |
| Epoprostenol, 7 to 14 days | 2,211,143 to 3,010,140 | 2,353,917 to 3,208,958 |
| Duration of training days | | |
| Treprostinil, 1 to 2 days | 2,610,242 to 2,608,894 | 2,781,438 to 2,779,690 |
| Epoprostenol, 14 to 21 days | 2,604,525 to 2,616,758 | 2,775,321 to 2,787,554 |
| Daily drug system coverage rate, $20 to $25 | 2,621,625 to 2,406,707 | 2,832,421 to 2,577,503 |
| ALOS for rehospitalizations per year, 5 to 55 days | 2,027,856 to 4,941,783 | 2,138,163 to 5,354,536 |
| Utilization of lost time, 8 h to 24 h | NA | 2,781,438 to 3,131,036 |
| Valuation of lost time, $17.18 to $34.36 per hour | NA | 2,781,438 to 2,952,234 |

*Savings are presented as cumulative values. ALOS Average length of stay; NA Not applicable*
hospitalizations due to adverse events associated with the route of administration and the epoprostenol delivery system.

The evidence provided by the present pharmacoeconomic evaluation suggests that using treprostinil to treat patients with PAH could provide cost savings from both the MoH and societal perspectives over a three-year period. The average savings per day was estimated to be $40 to $42 per patient.

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