Interferon-alfa in the treatment of patients with inoperable locally advanced or metastatic renal cell carcinoma: a systematic review

Christina Canil, MD, FRCPC,* Sebastien Hotte, MD, FRCPC;† Linda A. Mayhew, PhD;‡ Tricia S. Waldron;‡ Eric Winquist, MD, FRCPC§

Abstract

A systematic review was undertaken to determine whether interferon-alfa (IFN-α) is an effective treatment for patients with inoperable locally advanced or metastatic renal cell carcinoma (mRCC). MEDLINE, EMBASE, the Cochrane Library, guideline databases and relevant meeting proceedings were searched. Randomized clinical trials (RCTs) or meta-analyses comparing IFN-α-containing regimens to placebo or non-immunotherapy controls, and that reported response rate, survival, toxicity or quality of life data were eligible. Two systematic reviews and eight RCTs met the selection criteria. A Cochrane review updated in 2005 reported higher response rates and reduced one-year mortality based on 4 RCTs in patients who received IFN-α. Of the eight RCTs, three reporting objective response rate showed significant differences favouring IFN-α. Two of five trials reporting survival data showed longer median survival in the IFN-α group. Adverse effects of IFN-α were consistent across the trials with increased intensity and frequency concordant with increased IFN-α dose. Meta-analyses of seven RCTs for objective response and six RCTs for mortality favoured IFN-α: odds ratio 6.87 (95% Confidence Interval [CI], 3.29 to 14.35) and hazard ratio 0.79 (95% CI, 0.69 to 0.91), respectively. The effectiveness of IFN-α in mRCC has been subject to skepticism. As IFN-α has been used as a control arm in RCTs of new targeted therapies, information about its effectiveness remains relevant. These data confirm genuine, if modest, effectiveness of IFN-α in mRCC.

Résumé

Une revue systématique a été entreprise afin de déterminer si l’interféron alpha (IFN-α) représente un traitement efficace chez les patients atteints d’un hypernéphrome localement avancé ou métastatique et non opérable. Des recherches ont été effectuées dans les bases de données MEDLINE et EMBASE, dans la Bibliothèque Cochrane, les guides de pratique et les comptes rendus des réunions pertinentes. Les essais cliniques avec randomisation (ECR) ou les méta-analyses comparant des schémas contenant l’IFN-α à un placebo ou à un traitement témoin autre qu’une immunothérapie et qui faisaient état du taux de réponse, du taux de survie, des données sur la toxicité ou la qualité de vie pouvaient être inclus. Deux examens systématiques et huit ECR satisfaisaient aux critères de sélection. Une revue provenant de la Bibliothèque Cochrane et mise à jour en 2005 signalait des taux de réponses plus élevés et une mortalité réduite après un an en fonction de 4 ECR chez des patients ayant reçu de l’IFN-α. Sur les huit ECR retenus, trois faisant état du taux de réponse objective ont montré une différence significative en faveur de l’IFN-α. Sur les cinq essais faisant état de données sur la survie, deux ont montré une survie médiane plus longue dans le groupe sous IFN-α. Les effets indésirables de l’IFN-α étaient constants d’un essai à l’autre, l’intensité et la fréquence de ces effets variant en fonction de la dose d’IFN-α. Les méta-analyses de sept ECR pour dégager la réponse objective et de six ECR pour dégager les taux de mortalité étaient en faveur de l’IFN-α : rapport de cotes, 6.87 (intervalle de confiance [IC] à 95 %, 3.29 à 14.35) et rapport des risques instantanés, 0.79 (IC à 95 %, 0.69 à 0.91), respectivement. L’efficacité de l’IFN-α dans le traitement de l’hypernéphrome métastatique est parfois mise en doute. Comme l’IFN-α a été utilisé comme traitement témoin dans des ECR portant sur de nouveaux traitements ciblés, auxquels tous les patients n’ont pas nécessairement accès, les renseignements concernant son efficacité conservent toute leur pertinence. Ces données confirment l’efficacité réelle, quoique modeste, de l’IFN-α dans le traitement de l’hypernéphrome métastatique.

Introduction

In 2009, about 4600 new renal cell carcinoma (RCC) cases and 1600 RCC deaths were projected for Canada. Among patients with RCC at the time of their first diagnosis, 45% would present with localized disease, 25% would have locally advanced disease with lymph node or local organ involvement and the remaining 30% would present with metastases.

Patients who present with localized disease are best treated with surgery, but as many as 30% of these patients will eventually relapse. When patients present with or develop inoperable locally advanced or metastatic disease, the main intent of treatment is to effectively control symptoms and provide a chance of improved survival. Unfortunately, the treatment of late-stage RCC remains a challenge to oncologists and urologists; unlike other solid malignancies, advanced or metastatic RCC is highly resistant to most available chemotherapeutic agents.

Immunotherapy was first suggested as a treatment for advanced or metastatic RCC after occasional spontaneous tumour regressions, and the presence of anti-tumour immune responses were observed in patients with this neoplasm.
The major immunological approaches that have been investigated in these patients have included cytokines, either as single agents or in combination with other cytokines or chemotherapy. One of the first classes of cytokines to be evaluated was interferon. Interferons are naturally occurring glycoproteins that are produced in response to viral infections, antigens, and mitogens and are often induced by other cytokines like tumour necrosis factor (TNF) and interleukins. The anti-tumour activity of interferons is mediated by various mechanisms, such as immunomodulation, antiproliferative activity, inhibition of angiogenesis, regulation of differentiation, interaction with growth factors and modulation of gene expression. This systematic review assesses the effectiveness of interferon-alfa (IFN-α) for the treatment of advanced or metastatic RCC, based on the results of reported randomized controlled trials (RCTs).

Methods

Literature search strategy

Relevant RCTs, controlled clinical trials, meta-analyses, systematic reviews and practice guidelines were identified in electronic searches of MEDLINE (1966 through May 2009) and EMBASE (1980 through 2009 week 19).


Study selection criteria

Relevant articles and abstracts were selected by 4 reviewers. Papers were included if they were published RCTs, abstracts of RCTS, or meta-analyses that compared IFN-α-containing treatment regimens to regimens without IFN-α in patients with inoperable locally advanced or metastatic RCC. Comparison groups could include placebo, cytotoxic chemotherapy, hormonal therapies, such as medroxyprogesterone (MPA), and IFN-γ. Reports were required to provide data on at least one of the following outcomes: response rate, survival (overall, progression-free, and time-to-progression), toxicity and quality of life.

Studies that compared surgery or radiotherapy with IFN-α-containing treatment, compared IFN-α with angiogenesis inhibitors, or compared IFN-α with interleukin-2 (IL-2) were excluded, as these are examined in separate systematic reviews.

Synthesizing the evidence

For some eligible trials, odds ratios (OR) for overall mortality at 1 year and objective response and hazard ratios (HR) for overall mortality were available from a Cochrane meta-analysis by Coppin and colleagues. The analytic plan was to combine published data on these endpoints for all eligible trials, using meta-analysis. When the HR and its associated variance were available, those statistics were either extracted directly from the trial itself, from the Cochrane meta-analysis or were obtained through personal communication with trial authors. Otherwise, the HR was estimated indirectly from data extracted from published Kaplan-Meier curves using the methods of Parmar and colleagues. If data were not provided from which the HR could be derived or the authors did not provide the HR, the trial was not included in the meta-analysis. To estimate the overall effect of IFN-α, the data were combined using Review Manager version 4.2 (Cochrane Collaboration 2002, Review Manager, Oxford, England). Results are expressed as HR or OR with 95% confidence intervals (CI), where values <1.0 represent a benefit for IFN-α over the alternative (for HR and OR of mortality), and values >1.0 indicate a benefit for IFN-α (for OR of response). Use of a random effects model was planned.

Results

Literature search results

Two systematic reviews with meta-analyses and 8 RCTs met the inclusion criteria. No evidence-based guidelines were identified.

Systematic reviews with meta-analyses

The evidence from a meta-analysis published in 1999 was superseded by the results of a Cochrane review by Coppin and colleagues. Within the Cochrane review, the pooled results of 4 RCTs showed that IFN-α was associated with reduced 1-year mortality and greater remission rates compared with control therapy (MPA or vinblastine [VBL]). Remission, defined as the number of patients receiving a partial or complete response, was greater for IFN-α than control (12.5% vs. 1.5%) with a pooled OR of 7.61 (95% CI, 3.02–19.2). Interferon-α was also associated with reduced 1-year mortality (OR = 0.56; 95% CI, 0.40–0.77).

The 4 trials were also pooled using the methods of Parmar and colleagues to further explore the effect of IFN-α on mortality outcomes. The pooled overall HR for death was 0.74 (95% CI, 0.63–0.88), indicating a survival benefit for IFN-α.
over control therapy. The authors concluded that IFN-α demonstrated a modest improvement in remission rates and a consistent and statistically significant mortality reduction compared with a variety of controls. The analysis that compared trials using the recombinant subtypes IFN-α-2a and IFN-α-2b showed no evidence of statistical heterogeneity for either objective response or 1-year mortality.

Randomized controlled trials

Eight RCTs comparing IFN-α either alone or plus control therapy with control therapy alone met the inclusion criteria. The trials included 1360 patients, with patient accrual per trial arm ranging from 16 to 176. Patients were eligible for inclusion if they had histologically confirmed RCC and showed no signs of brain metastases. The median age of patients ranged from 56 to 63 years, and most were men (range, 59% to 75%) with good performance status (i.e., Eastern Cooperative Oncology Group or World Health Organization <2, Karnofsky >80% or Zubrod <2). The 8 RCTs provided a total of 17 comparisons. The RCTs reported outcome data for objective response rate, survival, disease progression and toxicity. No RCTs formally assessed quality of life.

Trial quality

Based on their reports, the quality of the trials was generally suboptimal. There were inconsistencies in data reporting in several of the trials. Two studies did not report the number of patients randomized per group. In addition, there were inconsistencies within a report regarding the number of patients reported, treated and randomized; in some cases, different publications of the same trial reported a different number of patients randomized. Some of these issues were resolved through personal correspondence either with the study authors or with the author of the Cochrane

<table>
<thead>
<tr>
<th>Table 1. Quality of eligible trials</th>
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<tbody>
<tr>
<td><strong>Trial</strong></td>
</tr>
<tr>
<td>Negrier 2005/2006/2007</td>
</tr>
<tr>
<td>Dutcher 2003</td>
</tr>
<tr>
<td>Hancock 2003</td>
</tr>
<tr>
<td>Pyrhönen 1999</td>
</tr>
<tr>
<td>Kriegmair 1995</td>
</tr>
<tr>
<td>Steineck 1990</td>
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<tr>
<td>Dexeus 1989</td>
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<tr>
<td>Foon 1988</td>
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</tbody>
</table>

*Trial stopped early on advice of independent data monitoring committee. NA = not applicable; NR = not reported.
### Table 2. Trial descriptions and outcomes: IFN-α containing regimens vs. control (8 RCTs, 8 comparisons)

<table>
<thead>
<tr>
<th>Trial</th>
<th>Treatment arms</th>
<th>Route, dose and schedule</th>
<th>No. patients randomized (evaluable)</th>
<th>Objective response rate %</th>
<th>Survival</th>
<th>Progression-free survival</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>OR</td>
<td>CR</td>
<td>PR</td>
</tr>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Negrier 2005/2006&lt;sup&gt;b&lt;/sup&gt;</td>
<td>IFN-α</td>
<td>sc 9 MU tiw</td>
<td>122 (115)</td>
<td>4.4&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.9&lt;sup&gt;d&lt;/sup&gt;</td>
<td>3.5&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>MPA</td>
<td>po 200 mg/d</td>
<td>123 (120)</td>
<td>2.5&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.8&lt;sup&gt;d&lt;/sup&gt;</td>
<td>1.7&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Dutcher, 2003&lt;sup&gt;f&lt;/sup&gt;</td>
<td>IFN-γ + IFN-α</td>
<td>iv 0.3 mg/m² daily x 5 d repeated q 3 wks sc 10 MU/m² daily</td>
<td>NR (49)</td>
<td>10</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>IFN-γ</td>
<td>sc 0.1 mg/m²/wk x 6 wks</td>
<td>NR (39)</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Hancock, 2003&lt;sup&gt;g,h&lt;/sup&gt;</td>
<td>IFN-α</td>
<td>sc 10 MU tiw x 12 wks</td>
<td>174 (167)</td>
<td>14&lt;sup&gt;i&lt;/sup&gt;</td>
<td>NR</td>
<td>NR</td>
</tr>
<tr>
<td></td>
<td>MPA</td>
<td>po 300 mg q d x 12 wks</td>
<td>176 (168)</td>
<td>1&lt;sup&gt;j&lt;/sup&gt;</td>
<td>NR</td>
<td>NR</td>
</tr>
<tr>
<td>Pythönen, 1999&lt;sup&gt;i&lt;/sup&gt;</td>
<td>IFN-α</td>
<td>iv 0.1 mg/kg q 3 wks</td>
<td>79 (79)</td>
<td>16.5</td>
<td>8.9</td>
<td>7.6</td>
</tr>
<tr>
<td></td>
<td>VBL</td>
<td>iv 0.1 mg/kg q 3 wks</td>
<td>81 (81)</td>
<td>2.5</td>
<td>1.2</td>
<td>1.2</td>
</tr>
<tr>
<td>Kriegmaier, 1995&lt;sup&gt;j&lt;/sup&gt;</td>
<td>IFN-α</td>
<td>sc 8 MU tiw&lt;sup&gt;k&lt;/sup&gt; iv 0.1 mg/kg q 3 wks</td>
<td>44 (41)</td>
<td>20</td>
<td>9</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>VBL</td>
<td>im 500 mg/wk</td>
<td>45 (35)</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Steineck, 1990&lt;sup&gt;k&lt;/sup&gt;</td>
<td>IFN-α</td>
<td>sc 10 MU/m² tiw&lt;sup&gt;l&lt;/sup&gt; dose was escalated wkl by 2.5 MU/m² to a max of 20 MU/m²</td>
<td>30 (NR)</td>
<td>6</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>MPA</td>
<td>im 1 g tiw x 5 wks; 1 g/wk</td>
<td>30 (NR)</td>
<td>3</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Dexeus, 1989&lt;sup&gt;l&lt;/sup&gt;</td>
<td>IFN-α</td>
<td>im 3 MU/m² d 1, 15; 5 MU/m² d 2,16, 10 MU/m² d 3-5,17-19</td>
<td>NR (16)</td>
<td>12</td>
<td>0</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>5-FU + CIS + DOX + MMC (FAMP)</td>
<td>iv 750 mg/m² d 1&lt;sup&gt;m&lt;/sup&gt;</td>
<td>NR (16)</td>
<td>12</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>Foon 1988&lt;sup&gt;n&lt;/sup&gt;</td>
<td>IFN-α</td>
<td>sc 2 MU/m² tiw</td>
<td>21(21)</td>
<td>2 PR</td>
<td>2 CR&lt;sup&gt;n&lt;/sup&gt;</td>
<td>NR</td>
</tr>
<tr>
<td></td>
<td>IFN-α</td>
<td>sc 1 MU/m² tiw</td>
<td>21(21)</td>
<td>2 PR</td>
<td>2 CR&lt;sup&gt;n&lt;/sup&gt;</td>
<td>NR</td>
</tr>
<tr>
<td></td>
<td>IFN-α + IFN-γ</td>
<td>47(47)</td>
<td>2 PR</td>
<td>2 CR&lt;sup&gt;n&lt;/sup&gt;</td>
<td>NR</td>
<td>NR</td>
</tr>
</tbody>
</table>

<sup>a</sup> Value α containing regimens vs. control (8 RCTs, 8 comparisons)

<sup>b</sup> Data extracted from survival curve. 

<sup>c</sup> Overall response rate is based on 153 patients and 156 patients in the IFN-α and MPA treatment groups at 12 weeks, respectively. The overall response rate at 6 months for the IFN-α and MPA was 8% and 1%, respectively. 

<sup>d</sup> Data was not used in the meta-analysis. 

<sup>e</sup> Only statistically significant differences are presented. 

<sup>f</sup> No. patients in each of the IFN-α and MPA treatment groups at 12 weeks, respectively. The overall response rate at 6 months for the IFN-α and MPA was 8% and 1%, respectively. 

<sup>g</sup> Ten people received purified, 

<sup>h</sup>Two responses occurred in the combination arm. 

<sup>i</sup> Ten people received purified, 

<sup>j</sup>Data was not used in the meta-analysis. 

<sup>k</sup>Data was not used in the meta-analysis. 

<sup>l</sup>Data was not used in the meta-analysis. 

<sup>m</sup>Data was not used in the meta-analysis. 

<sup>n</sup>Data was not used in the meta-analysis. 

<sup>o</sup>Data was not used in the meta-analysis. 

<sup>p</sup>Data was not used in the meta-analysis. 

<sup>q</sup>Data was not used in the meta-analysis. 

<sup>r</sup>Data was not used in the meta-analysis. 

<sup>s</sup>Data was not used in the meta-analysis. 

<sup>t</sup>Data was not used in the meta-analysis. 

<sup>u</sup>Data was not used in the meta-analysis. 

<sup>v</sup>Data was not used in the meta-analysis. 

<sup>w</sup>Data was not used in the meta-analysis. 

<sup>x</sup>Data was not used in the meta-analysis. 

<sup>y</sup>Data was not used in the meta-analysis. 

<sup>z</sup>Data was not used in the meta-analysis. 

<sup>α</sup> α = interferon-alfa; IFN-α = interferon-gamma; im = intramuscularly; IU = International Units; iv = intravenous; kg = kilogram; m = meters; mg = milligrams; MMC = mitomycin C; MPA = medroxyprogesterone acetate; mos = months; MU = Million Units; NR = not reported; NS = non-significant; OR = objective response; po = per oral; PR = partial response; q = every; RC Ts = randomized controlled trials; sc = subcutaneous; tiw = three times a week; VBL = vinblastine; vs. = versus; wk(s) = week(s); yr = year. 

Values are given as median (mos) for survival analysis.
review. However, the new data provided often conflicted with the data presented in the original papers. Furthermore, 1 study reported 2 responses in the combination arm of the trial and one in each of the other arms but did not report the type of response. Two studies did not report mortality data. Study quality elements of the included trials are summarized in Table 1.

**Outcomes**

Outcome data from the RCTs for objective response rate, survival and disease progression are reported in Table 2. All 3 RCTs reporting objective response rate showed a statistically significant difference in favour of IFN-α. Hancock and colleagues reported longer median survival for patients receiving IFN-α than MPA (9 vs. 6.75 months, p = 0.013). Pyrhönen and colleagues reported that patients receiving IFN-α-2a combined with VBL had a longer median survival than that of patients receiving VBL alone (15.5 vs. 8.7 months, p = 0.0049). Of the 3 remaining trials, 1 found no difference between trial arms, and 2 did not provide statistical comparisons. A sixth trial reported no difference in median survival between groups, but did not provide data. One-year survival data were reported in 2 trials and extracted from survival curves in 4 trials. Two trials did not report survival outcomes.

Disease progression was assessed in 3 trials. The trial by Pyrhönen and colleagues reported that IFN-α-2a in combination with VBL demonstrated significantly longer progression-free survival than did treatment with VBL alone (3 vs. 2.1 months, p = 0.0001).

The adverse effects of IFN-α were reasonably consistent from trial to trial, although they increased in intensity and frequency with increased IFN-α dose. Reports of the toxicities from a large RCT as representative of IFN-α toxicity showed increased rates of lack of appetite (51%), tiredness (68%), nausea/vomiting (26%/9%), lack of energy (65%), dry mouth (41%), shivering (23%) and depressed mood (25%) with IFN-α after 4 weeks of treatment. Increased rates of lack of appetite, tiredness, lack of energy, dry mouth and shivering persisted at 12 weeks. Other reported symptoms included irritability, worrying, sore muscles, general pain, nervousness, despondent or tense feelings, difficulty sleeping, headaches, dizziness, decreased sexual interest, restlessness, anxiety, constipation, diarrhea, tingling in hands and feet, difficulty concentrating, sour mouth, loss of hair, shortness of breath, hoarseness and burning eyes.

**Meta-analysis**

Overall response and mortality were considered the primary endpoints for the meta-analysis. Because of the previously mentioned issues with data consistency, study authors and the author of the Cochrane review were contacted to determine the data on which to base results. Responses from the authors were used in some cases as opposed to the data reported in trial reports for the meta-analysis.

The response data from 7 trials were pooled in a meta-analysis. The number of patients randomized to each treatment arm was not available for 1 trial; therefore, this trial was not included in the meta-analysis. The results of the meta-analysis appear in Fig. 1. The meta-analysis of the 7 trials produced an OR of 6.87 (95% CI, 3.29-14.35; p < 0.00001).
Mortality data suitable for meta-analysis were reported in 6 trials and combined in a meta-analysis. The results of the meta-analysis appear in Fig. 2. The HR for mortality after treatment with IFN-α was 0.79 (95% CI, 0.69-0.91; p = 0.001).

**Discussion**

This review identified 8 RCTs that directly evaluated the use of IFN-α in locally advanced or metastatic RCC. These trials compared IFN-α alone or in combination with control therapies considered to have little or no activity in RCC. In our meta-analysis of 6 trials, the overall HR for death was 0.79, indicating a 21% reduction in the risk of death for patients treated with IFN-α over the time periods of follow-up of the RCTs. Toxicity was higher with IFN-α therapy and, unfortunately, health-related quality of life was not evaluated. However, the odds of objective response were almost 7 times higher for patients receiving IFN-α-containing regimens (4.4% to 20%) compared with patients in control groups (0 to 3%). Heterogeneity was minimal in these analyses. None of the included RCTs used placebo control groups, which is an inherent limitation of this data set. This could potentially exaggerate survival differences between IFN-α and control due to the detrimental effects of control therapy, a development that was more likely to occur when chemotherapy was the control. Medroxyprogesterone acetate and IFN-γ have not shown detrimental effects on survival, and analysis of these trials alone shows similar pooled survival results (data not shown). Only 1 trial included in the analyses used chemotherapy (single-agent VBL) as control. This trial did have the most extreme HR for overall mortality, favouring IFN-α therapy. However, VBL has a low objective response rate in RCC (7%), is considered a mild cytotoxic drug, and was also included in combination with IFN-α in the experimental arm of this trial. A sensitivity analysis excluding the trial provided an overall HR for death of 0.83 (95% CI, 0.71-0.96, p = 0.01). The limitations of published data meta-analysis have been well-described and are potentially applicable to these results; nevertheless, we consider the results a comprehensive and robust synthesis of the best currently available clinical data.

Overall, toxicity appeared worse with IFN-α compared with non-IFN-α therapy. Toxicity for IFN-α is well-known and consistent from trial to trial; therefore, we presented toxicity data from a large randomized trial that reported increased rates of lack of appetite, tiredness, lack of energy, dry mouth and shivering with IFN-α after 12 weeks of treatment. No toxic deaths were reported; however, these data were not reported in most of the studies. The general opinion is that IFN-α regimens are associated with substantial toxicity; the magnitude of this toxicity may be underestimated in clinical trials due to patient selection factors, such as performance status and under-reporting.

Doses and modality of administration of IFN-α differed across the trials. In 5 trials, IFN-α was administered subcutaneously at doses ranging from 2 to 10 MU/m² on a thrice-weekly schedule. In 2 trials, IFN-α was administered intramuscularly at doses ranging from 3 to 10 MU/m², and in another it was administered either subcutaneously or intramuscularly starting at 3 MU and increasing to 18 MU. Whether there is a dose response to IFN-α is unclear; however, it is likely that toxicity is dependent on dose and schedule. In addition, there is no evidence of a difference in efficacy between recombinant IFN-α-2a and IFN-α-2b, or clear evidence of benefit of adding chemotherapy to IFN-α. In view of this, the consensus of the authors was that it is reasonable to use the dose and schedule from the largest RCT showing benefit.

This trial gave an initial
tial dose of 5 MU subcutaneously followed by 10 MU subcutaneously on a thrice-weekly schedule for a total of 12 weeks, unless progressive disease or objective response was seen. Treatment could be continued after 12 weeks in responding patients. In view of these data and the toxicities of IFN-α, the value of treatment beyond 12 weeks in non-responding patients is questionable and should be considered on an individual basis.

Despite many years of research, the prognosis for patients with inoperable locally advanced or metastatic RCC had not changed until recently, and very few therapeutic options existed for these patients. Our synthesis of the data from randomized trials of IFN-α-based immunotherapy confirms that IFN-α has anti-tumour activity in RCC, provides a genuine if modest survival benefit in this patient population, and should be considered as a potential treatment option. Evidence from randomized trials of angiogenesis inhibitors (i.e., sunitinib, sorafenib and temsirolimus) show that these agents are of superior clinical effectiveness to IFN-α, with acceptable toxicity. The clinical benefits observed with these agents make them the preferred treatment modality. In particular, the low objective response rate (ORR) seen with IFN-α (6% to 20%) suggests that drugs, such as sunitinib (ORR=33% to 40%), may be preferred in patients with disease involving critical organs where prompt disease shrinkage to secure survival may be necessary.

As not all patients may have access to the newer angiogenesis therapies due to their costs, information about the effectiveness of IFN-α is still of value. Despite IFN-α and recent advances with other new drugs, patients with inoperable locally advanced or metastatic RCC continue to have an incurable malignancy, and further research to improve disease control and cure is necessary.

Conclusion

Until recently, very few systemic therapeutic options existed for patients with inoperable locally advanced or metastatic RCC. Immunotherapy with IFN-α can be considered a treatment option to modestly improve survival and disease control in this patient population. However, given the toxicity profile of IFN-α, patient factors, such as age and performance status, must be taken into consideration and may affect patients’ ability to tolerate therapy and benefit from it. Further, angiogenesis inhibitors have expanded the treatment repertoire for RCC and appear to have superior effectiveness compared to IFN-α. In view of this, the role of IFN-α in the treatment of RCC is less clear. However, as not all patients may have access to the newer therapies due to the cost of these therapies, information about the effectiveness of IFN-α is still of value.

Locally advanced or metastatic RCC remains an incurable disease, current treatments remain palliative, and further research is warranted. Whenever possible, patients should be encouraged to participate in clinical trials.

Competing interests: The members of the Genitourinary Cancer Disease Site Group disclosed potential conflicts of interest relating to this systematic review. One author (SH) reported grant/research support from two companies with competing treatments, and serving on an advisory board for one company with a competing treatment. One author (CC) reported serving on an advisory board for two companies with competing treatments. No further conflicts were declared by the authors.

Acknowledgements: The Genitourinary Cancer Disease Site Group would like to thank Dr. Christina Camill, Dr. Sebastian Hotte and Dr. Eric Winquist for taking the lead in drafting this systematic review with meta-analyses. The authors would also like to thank Dr. Janice Dutcher, Ms. Judi Manola, and Ms. Sylvia Negier for their personal correspondence clarifying data, Dr. Chris Coppin for his correspondence concerning his review, Ms. Kate Bak for contributions to an early version of this report, and Ms. Cindy Walker-Bilks for assistance in preparing this manuscript.

This paper has been peer-reviewed.

References


**Correspondence:** Dr. Christina Canil, The Ottawa Hospital Regional Cancer Centre, 503 Smyth Rd., Ottawa, ON K1H 1C4; fax: 613-247-3511; ccanil@ottawahospital.on.ca