Original article

Description of the EuroTARGET cohort: A European collaborative project on TArgeted therapy in renal cell cancer—GEnetic- and tumor-related biomarkers for response and toxicity

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Abstract

Objective: For patients with metastatic renal cell cancer (mRCC), treatment choice is mainly based on clinical parameters. With many treatments available and the limited response to treatment and associated toxicities, there is much interest in identifying better biomarkers for personalized treatment. EuroTARGET aims to identify and characterize host- and tumor-related biomarkers for prediction of response to tyrosine kinase inhibitor therapy in mRCC. Here, we describe the EuroTARGET mRCC patient cohort.

Methods and materials: EuroTARGET is a European collaborative project designed as an observational study for which patients with mRCC were recruited prospectively in 62 centers. In addition, 462 patients with mRCC from previous studies were included. Detailed clinical information (baseline and follow-up) from all patients was entered in web-based case record forms. Blood was collected for germline DNA and pharmacokinetic/pharmacodynamic analyses and, where available, fresh-frozen tumor material was collected to perform tumor DNA, RNA, kinase, and methylome analyses.

Results: In total, 1,210 patients with mRCC were included. Of these, 920 received a tyrosine kinase inhibitor as first-line targeted treatment (sunitinib [N = 713, 78%], sorafenib [N = 41, 4%], or pazopanib [N = 166, 18%]) and had at least 6 months of outcome assessment (median follow-up 15.3 months [interquartile range: 8.5–30.2 months]). Germline DNA samples were available from 824 of these patients, fresh-frozen tumor material from 142 patients, fresh-frozen normal kidney tissue from 95 patients, and tissue microarrays created from formalin-fixed paraffin-embedded tumor material from 247 patients. Of the 920 patients, germline DNA variant chip data were successfully generated for 811 patients (Illumina HumanOmniExpress BeadChip). For 80 patients, next-generation exome sequencing of germline and tumor DNA was performed, tumor RNA sequencing was performed for 124 patients, kinase activity measured and processed for 121 patients (PamChip), and methylome data (Illumina Infinium HumanMethylation450 BeadChip) were created for 116 RCC tissues (and 23 normal kidney tissues). For 73 out of the 920 patients, all platform data types were generated. In addition, 40 patients were included in a pharmacokinetic/pharmacodynamic phase IV substudy.

Conclusions: Analysis of EuroTARGET cohort data will contribute to personalization of therapy for patients with mRCC. The extensive clinical data and multiplatform EuroTARGET data will be freely available. © 2017 The Authors. Published by Elsevier Inc. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

Keywords: Metastatic renal cell carcinoma; Therapy response; Tyrosine kinase inhibitor; Biomarker; Transcriptomics; Genomics

1. Introduction

With more than 121,000 newly diagnosed patients and 52,000 deaths each year, kidney cancer is the seventh most common cancer in Europe [1]. Further, 90% of all kidney cancers are renal-cell carcinomas (RCC). The prognosis of RCC is highly dependent on stage. Surgery is effective for the 70% to 80% of patients with localized disease, leading to 5-year relative survival rates of more than 70% [2]. However, ~25% of patients have metastatic RCC (mRCC) at first diagnosis, and ~25% of patients with localized disease develop metastases after surgery [3,4].

Until the arrival of tyrosine kinase inhibitors (TKIs), treatment options in mRCC were limited, and 5-year relative survival was only 5% to 10%. Randomized clinical trials showed that TKI agents, directly targeting tumorigenic and angiogenic pathways (reviewed in [5]), significantly improved the outcomes of these patients [6]. Several first-line TKI treatment options are now available, such as sunitinib, pazopanib, or bevacizumab plus interferon-alpha [6]. Most patients experience disease stabilization or response for a median of ~12 months [7]. However, TKI treatment is extremely expensive (sunitinib was estimated in the UK to cost £71,462 per quality adjusted life year gained [8]), 15% to 20% of patients experience immediate disease progression despite treatment [7], nearly all patients eventually become resistant, and toxicity is common and leads to dose reduction. In addition, first-line treatment options for patients with mRCC will likely increase in coming years (including immune checkpoint inhibition) [9]. There is, hence, much interest in tools for prediction of individual therapy response and acquired resistance to TKIs to optimize treatment outcome while reducing unnecessary drug use and expenses, and improving human health and quality of life.

Currently, treatment choice in mRCC is based on risk grouping of patients by clinical parameters such as the patient’s performance status, and serum biochemical measurements, and histological features of the tumor [10]. A comparison study into several clinical risk grouping models, including that of the International Metastastic Renal Cell Carcinoma Database Consortium [11] and the Memorial Sloan Kettering Cancer Center (MSKCC) [12], showed modest discriminatory values for survival (area under the receiver operating curve ~0.66) and indicated that addition of tumor-specific or patient-specific biomarkers is likely required for the improvement of the accuracy of these models [13].

Advances in high-throughput technologies have paved the way to personalized medicine using biomarkers. For mRCC, potential prognostic molecular biomarkers such as PBRM1, BAP1, and KDM5C tumor mutations [14]; IL8 [15], VEGF, and PI GF levels [16]; ABCB1 and VEGFR-3 germline polymorphisms [17]; and miRNA levels have been identified [18,19]. However, there are no validated biomarkers yet that can guide personalization of therapy in patients with mRCC.

In this framework EuroTARGET was initiated, a “European collaborative project on TArgeted therapy in Renal cell cancer: GEnetic and Tumor-related biomarkers for response and toxicity.” The overarching goal of this
multicenter observational study is to identify and characterize host and tumor-related biomarkers that can be used to distinguish expected responders from nonresponders for targeted therapy in mRCC. Here, we describe the cohort of recruited patients, their characteristics, and the collected data.

2. Material and methods

2.1. Study design and patient recruitment

EuroTARGET is an international, multicenter observational study that started in March 2011. The study was approved by the ethics committee at each participating center. A total of 748 patients were recruited prospectively in 62 centers; 36 in The Netherlands, 16 in Spain, 8 in Germany, 1 in Romania, and 1 in the United Kingdom. Patient identification and invitation procedures differed per country (Supplementary material), but at all locations, the following were the inclusion criteria: patients gave written informed consent, were at least 18 years of age, and had newly diagnosed metastatic renal-cell carcinoma. Patients were enrolled in the study regardless of RCC subtype and (pre)treatment.

In addition to the prospectively recruited patients, 462 patients with mRCC from previous studies (“historical patients”) were included in EuroTARGET. Of these, 56 patients were retrospectively included in PERCEPTION, a Dutch population-based study with patients diagnosed between January 2008 and December 2010 [20]; 89 patients were enrolled between October 2007 and December 2010 by the Spanish Oncology Genitourinary Group [21]; 153 patients were included between June 2004 and October 2010 by a Dutch working group of 6 university hospitals focussing on sunitinib-induced toxicity [22]; and 35 patients were recruited to a study on the epidemiology and inheritance of renal cancer conducted by deCODE genetics in Iceland since 2001 [23]. The Radboud university medical center (Radboudumc) included another 66 patients, collected in a prospective biobank of patients diagnosed with cancer. The Central European Society for Anticancer Drug Research included an extra 63 patients, collected in prospective biobanks from Saarland University Medical Center in Homburg, Germany, Jena University Hospital in Germany, and University Hospital Graz in Austria. All biobanks were approved by the local ethics committees (Supplementary material).

2.2. Collection of clinical data

Clinical information from all patients was collected by medical file review and entered in web-based case record forms (CRFs) (Supplementary material and Table 1). All data were managed, exchanged cross-border, and used according to the data protection laws in Europe. Data included demographic information, baseline clinical characteristics, lab values at start of treatment of metastasis, number of locations and location of metastases, clinical data at start of treatment of metastasis, comorbidities at start of treatment of metastasis, and final information on patient.

2.3. Collection of biomaterials

Blood samples were collected from almost all patients for germline DNA isolation. In 12 German and Dutch centers, up to 12 plasma samples were collected per patient before and during sunitinib or pazopanib treatment, for measurement of drug and metabolite concentrations. In addition, tumor material from the kidney and normal kidney tissue was collected during nephrectomy if possible, and freshly frozen. This fresh-frozen material was mainly

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Table 1

<table>
<thead>
<tr>
<th>Clinical information available in EuroTARGET web-based case record forms</th>
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<tbody>
<tr>
<td>Patient characteristics</td>
</tr>
<tr>
<td>Date of birth</td>
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<tr>
<td>Sex</td>
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<tr>
<td>Eligibility criteria</td>
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<tr>
<td>Date of diagnosis metastasis</td>
</tr>
<tr>
<td>General information at diagnosis</td>
</tr>
<tr>
<td>Height</td>
</tr>
<tr>
<td>Bilateral renal cell carcinoma</td>
</tr>
<tr>
<td>Tumor assessment at start of treatment</td>
</tr>
<tr>
<td>cTNM classification</td>
</tr>
<tr>
<td>pTNM classification</td>
</tr>
<tr>
<td>Comorbidities at start of treatment of metastasis</td>
</tr>
<tr>
<td>Treatment line</td>
</tr>
<tr>
<td>Clinical data at start of treatment of metastasis</td>
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<tr>
<td>Number of locations and location of metastases</td>
</tr>
<tr>
<td>Lab values at start of treatment of metastasis</td>
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<tr>
<td>Drug treatment</td>
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<tr>
<td>Dosing schemes</td>
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<tr>
<td>(Concomitant) nondrug treatment—surgery/radiotherapy</td>
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<tr>
<td>Toxicities</td>
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<tr>
<td>Responses</td>
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<tr>
<td>New lesions</td>
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<tr>
<td>Final information on patient</td>
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<tr>
<td>Date of death</td>
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<td>Last registration date</td>
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</table>

\[\text{cTNM} = \text{clinical tumor-nodes-metasasis}; \text{pTNM} = \text{pathological tumor-nodes-metasasis} \]
collected in the German and Dutch academic centers. Paraffin-embedded tumor material from the kidney was not collected specifically for EuroTARGET, but it was collected from patients with histologically confirmed RCC at the local pathology departments. Slides were collected for central pathology review of tumor subtype and histopathological features by one of the four expert uropathologists from Spain, The Netherlands, Germany, and the United Kingdom. In The Netherlands, paraffin-embedded material was also used to construct tissue microarrays (TMAs) containing three 3 mm cores from representative tumor areas per patient. Blood or DNA samples or both and freshly frozen samples were coded and stored at the central EuroTARGET biobank at the Radboudumc, The Netherlands.

2.4. Platform and pharmacokinetic/pharmacodynamic analyses

EuroTARGET encompasses multiplatform omics profiling and pharmacokinetic/pharmacodynamic (PK/PD) analyses. Genome-wide germline DNA variation data were measured using Illumina HumanOmniExpress BeadChips. Tumor material was profiled using next-generation whole-exome sequencing (Illumina; tumor DNA and matched germline DNA), RNA sequencing (Illumina), PamChip kinase assays, and Illumina Infinium HumanMethylation450 BeadChips. PK/PD models were developed using NONMEM 7.3 software.

2.5. Availability of data

All clinical and platform data generated in EuroTARGET will be made freely available in an anonymized way for the research community as of March 2018. The data can be accessed through the European Genome-phenome Archive (EGA) which is the controlled access repository under the European Bioinformatics Institute (EMBL-EBI). Interested parties will be able to find the EuroTARGET project under the Studies section.

3. Results

In total, 1,210 patients with mRCC were included in EuroTARGET, of which 748 were collected prospectively and 462 were available from historical (prospective) series at the start of EuroTARGET. Of the 1,210 patients, we selected the 979 patients (81%) who received sunitinib, sorafenib, or pazopanib as first-line TKI (remainder of patients did, for example, have no treatment or were treated with an mTOR inhibitor or other TKI). Prior cytokine therapy was allowed. To enable informative future analyses, we only focus on the subset of 920 patients for whom outcome could be assessed for at least 6 months (24 weeks) (Fig. 1).

Table 2 displays the origin of the 1,210 patients collected for EuroTARGET and of the 920 patients who were selected as being relevant for future analyses.

3.1. Baseline characteristics

Baseline characteristics of the total patient population stratified by first-line TKI are shown in Table 3 (see Tables S1a and S1b in supplementary material for baseline characteristics.

Table 2

<table>
<thead>
<tr>
<th></th>
<th>All collected patients</th>
<th>Patients to be included in analyses</th>
</tr>
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<tbody>
<tr>
<td>Prospective recruitment</td>
<td>N = 1,210</td>
<td>N = 920</td>
</tr>
<tr>
<td>Patients from historical series</td>
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<thead>
<tr>
<th>Country</th>
<th>EuroTARGET</th>
<th>PERCEPTION</th>
<th>SUTOX</th>
<th>Radboudumc</th>
<th>Spain</th>
<th>EuroTARGET</th>
<th>SOGUG</th>
<th>United Kingdom</th>
<th>EuroTARGET</th>
<th>Romania</th>
<th>EuroTARGET</th>
<th>Germany</th>
<th>EuroTARGET</th>
<th>CESAR</th>
<th>Austria</th>
<th>CESAR</th>
<th>Iceland</th>
<th>DeCODE</th>
<th>Total</th>
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<tr>
<td>The Netherlands</td>
<td>264</td>
<td>56</td>
<td>150</td>
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<td>187</td>
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<td>169</td>
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<tr>
<td>United Kingdom</td>
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<td>Germany</td>
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<td>Austria</td>
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<td>Iceland</td>
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CESAR = Central European Society for Anticancer Drug Research; PERCEPTION = Dutch population-based registry of mRCC patients; SOGUG = Spanish Oncology Genitourinary Group; SUTOX = Dutch working group of six university hospitals focusing on sunitinib-induced toxicity.
3.2. Follow-up data

Patient follow-up was completed up to March 1, 2016, and censored at this date if patients were still alive, which was the case for 241 patients (26%). A total of 478 patients (52%) were followed until their death. For 39 patients (4%), date of death was known, but follow-up was not completed up to the date of death. For another 162 patients (18%), follow-up information was not complete up to March 1, 2016, nor was date of death known. These patients were censored at the last date registered in the CRF. Median follow-up time from start of TKI treatment until death or censoring for all patients was 15.3 months (interquartile range [IQR]: 8.5–30.2 months).

Progression is defined as relapse or progressive disease, development of a new lesion, or death. We calculated PFS time as the time between start of TKI treatment and date of progression, death, or censoring (whichever happened first), regardless of duration of TKI treatment. OS time was calculated as the time between start of TKI treatment and date of death or censoring. A total of 681 patients (74%) experienced a progression event and 517 (56%) died during follow-up. Median PFS time was 10.8 months (IQR: 4.4–24.6 months) and median OS was 26.1 months (IQR: 10.6–51.3 months) (Fig. 2).
We also evaluated PFS for the time period in which patients were actually using TKI treatment (sunitinib, sorafenib, or pazopanib). In this scenario, patients were censored when they stopped TKI treatment, and PFS time was calculated as the time between start of TKI treatment and progression, death, or censoring (whichever came first). With this definition, 546 patients (59%) experienced a progression event, and median PFS time was 11.7 months (IQR: 5.0–30.1 months).

Table 4 shows the follow-up characteristics of the total patient population and separately for the different first-line TKIs used (see Supplementary Tables S3a and S3b for follow-up characteristics for the prospectively recruited and historical patients).

### 3.3. Biomaterial and platform data

Germline DNA was available from 824 of 920 patients and fresh-frozen tissue from 145 patients (primary kidney tumor, \(N = 142\); normal kidney, \(N = 95\)). TMA were created from 247 patients. Platform data were successfully generated for the following number of patients (out of the 920): germline DNA variation chip data for 811, whole-exome sequencing data (tumor and germline DNA) for 80, RNAseq data (tumor mRNA and miRNA) for 124, processed kinase activity data for 121, and methylation data for 116 RCC tissues (and 23 normal kidney tissues). For 73 patients, all platform data are available. PK/PD data are available from 40 patients.

### 4. Discussion

EuroTARGET is a European collaborative project, which aims to discover and validate biomarkers to personalize treatment of patients with mRCC. Here, we describe the EuroTARGET patient population, a large, extensively phenotyped cohort of 920 patients with mRCC. For most patients, genome-wide germline DNA variation data are available, making this the largest cohort for prognostic
germline genetic biomarker studies in sunitinib-treated patients with mRCC. For 73 patients, germline genome, tumor genome, transcriptome, kinome activity, as well as methylome data are available, allowing for an integrated analysis of multiplatform data.

In recent years, a number of renal cancer biorepositories with extended platform data have been generated. For example, the International Cancer Genome Consortium (https://icgc.org/), including, among others, The Cancer Genome Atlas Project [24], has profiled more than 1,200 patients with renal cancer at the DNA, RNA, protein, and epigenetics level. These data have been very valuable for insight into the existence of molecular subtypes [25]. However, the number of patients with mRCC and clinical data is limited, restricting the value of these data for prognostic biomarker studies in mRCC. Indeed, potentially relevant prognostic biomarkers for mRCC have, to date, mainly been derived from (randomized) clinical trials [14,18].

In contrast to clinical trial biorepositories with strict inclusion and exclusion criteria, EuroTARGET is of an observational nature. Therefore, patients should be more reflective of the general TKI population and results better generalizable. However, it also has disadvantages such as the dependence on information that is registered in medical files for information retrieval. Also, more than 60 centers and 5 European countries were involved in patient recruitment, resulting in a number of challenges. For example, the start of patient recruitment was severely delayed because of difficulties in obtaining ethical approval for this observational study that was erroneously regarded as a clinical trial by many of the recruitment centers.

Inclusion of patients from historical series in EuroTARGET substantially increased the number of available patients. It also poses some concerns, as these patients were sampled for projects with different aims, in different time periods, and using different inclusion procedures. For instance, we observed a median PFS and OS time of 8.9 and 20.7 months in the historical patient series compared to 11.8 and 30.3 months in the prospectively recruited patients, possibly reflecting improvement in mRCC treatment over time. Although biomarkers can be identified regardless of this, the quantitative effect estimate of the biomarker may not be representative for all current patients with mRCC.

The importance of replication and validation of biomarker findings has been stressed in many publications [26]. The original objective of EuroTARGET was to perform a two-stage inclusion of patients and samples to allow for both a discovery and a replication cohort within the consortium. However, owing to recruitment difficulties, the distinction between discovery and replication cohort has been abandoned. Instead, we will use external mRCC patient cohorts for replication [27–29]. Also note that functional validation of biomarker findings using, for example, in vitro studies in established RCC cell lines are integral parts of EuroTARGET.

Currently, analyses of clinical and platform data (separately and integrated), PK/PD analyses, and functional studies are ongoing within the EuroTARGET consortium. We hereby hope to improve understanding of the critical molecular and resistance pathways involved in TKI therapy and to define new validated risk stratification criteria to be used in personalized mRCC patient management.

5. Conclusions

EuroTARGET is a European collaborative project including 920 patients with mRCC treated with sunitinib, sorafenib, or pazopanib. EuroTARGET data will be freely available from March 1, 2018. We hope that easy access will promote the uptake of EuroTARGET data by the research community, and thereby the progress in personalization of therapy for patients with mRCC.

Acknowledgments

We would like to thank all patients who participated and their treating physicians for inviting them.

Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at http://dx.doi.org/10.1016/j.urolonc.2017.03.009.

References


