Nimotuzumab in Advanced Cervical Cancer: Safety and Efficacy Profile

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Abstract

The EGF receptor is over expressed in tumors of epithelial origin and is associated with a poor prognosis of the disease. An Expanded Clinical Access Program was carried out, in which patients with cervical uterine cancer in advanced stages of the disease without another therapeutic alternative were treated with nimotuzumab, anti-epidermal growth factor receptor humanized monoclonal antibody to evaluate the safety and efficacy of this therapy. Seventy-five patients with confirmed histological diagnosis of advanced cervical cancer were included, with previous onco-specific treatments (surgery, radiotherapy or radiochemotherapy), with ECOG at the inclusion ≤ 3 and life expectancy of more than three months. The mAb was administered in two stages: The induction stage with weekly administration, once a week in a dose of 200 mg intravenously; for twelve weeks and the second or maintenance stage with the same dose but every 14 days, until unmanageable toxicity or the decision of the patients. The therapy was well tolerated with an acceptable safety profile, not exceeding 2% the severe adverse events related to nimotuzumab treatment. The median overall survival exceeded the expectation by arriving at 23.5 months. We suggest follow up to a clinical trial in more early stage of the disease to combine nimotuzumab with chemoradiotherapy.

Keywords

Nimotuzumab, Cervical cancer, Advanced, General survival, Safety

Introduction

Cervical cancer, despite being one of the preventable locations with simple diagnostic methods such as the Papanicolaou test and more recently with the detection of DNA (or RNA) of HPV virus types 16 and 18 in cervical cells [1], it is maintained with an overall incidence of 528,000 cases in 2012 with an annual increase of 266,000 new cases. Eighty-five percent of the cases are diagnosed in developing countries [2]. Approximately 12,990 new cases were diagnosed in the US in 2016 and 4120 women died of this cause. Although the disease has decreased due to screening and preventive vaccination programs against the most oncogenic subtypes 16 and 18 of Human Papillomavirus (HPV), the incidence remains high in women of Latin, black and Asian origin [3].

According to PAHO in Latin America and the Caribbean, cervical cancer mortality is 3 times higher than in the US and Canada, with around 92,132 cases and 37,640 deaths annually, reporting a direct association with socioeconomic development and access to Health programs [4]. In Cuba, according to the Statistical Yearbook of Health, it is the 5th cause of mortality with 512 cases in 2016, with the highest number of deaths between 40 and 59-years-old [5-7].

It is estimated that 35% of cervical tumors are diagnosed as a locally advanced disease (stage Ib2-IIb according to the staging system of the International Federation of Gynecology and Obstetrics (FIGO)) with an estimated survival rate of 5 years in 60% of the cases, using the standard treatments of concurrent chemoradiotherapy based on platinum [8-10]. One percent
of these patients become as persistence (around 20%) or in relapse of the disease, where the platinum-based chemotherapy (Carboplatin and Cisplatin), the Taxans (Paclitaxel and Docetaxel), the Gemcitabine and the Topotecan are presented as the main therapeutic option [11].

Passive immunotherapy associated with the epidermal growth factor receptor (EGF R), using monoclonal antibodies (MAbs) have been demonstrate the specificity recognition of the receptor by the antibody and the inhibition of the ligands binding to it, evidencing an inhibitory effect of the mytogenic stimulation of the malignant cells [12].

In immune-histochemistry evaluations, it was found that over expression of EGF-R in squamous cells carcinoma of the uterine cervix has a direct relationship with the rate of tumor growth [13]. This receptor is associated with the poor prognosis of the disease because participate in the processes of proliferation of malignant cells, the invasiveness of healthy tissues, in addition to favoring the mechanisms of angiogenesis and inhibiting apoptosis [14].

The humanized Mab nimotuzumab (IgG1) recognizes the domain III of the EGF-R with a $10^{-9}$ mol/L constant of dissociation (considered as intermedia in relation to those reported for this type of antibodies) [15]. It avoids the interaction of EGF, (and other ligands) to its receptor and the consequent signals transduction to the nucleus, which favors blocking all its activating functions [15].

Safety and efficacy of nimotuzumab as a single agent or in combination with radio and chemotherapy in tumors from head and neck, glioma, esophagus, pancreas and more recently in advanced persistent or recurrent cervical cancer has been reported; showing an excellent safety profile, control of antitumor activity and improvement in survival [16-18].

Through the present, it is evaluated the safety and efficacy of nimotuzumab in patients with cervical cancer in advanced disease (relapse, tumor persistence or metastatic and refractory to previous oncospecific treatments), without therapeutic alternatives at the time of being included in the study.

Methods

A multicenter, prospective Expanded Clinical Access Program was performed. It was approved by the Institutional Ethics Committees for Clinical Research (six Cubans hospitals) and officially communicated to the Cuban Drug Regulatory Agency (CECMED).

Seventy-five patients were included since 2008 to 2010. All of them fulfilled the follow inclusion criteria:

- Informed consent to participate in the research duly signed, confirmed cito-histological diagnosis of cervical cancer, persistent or relapsing or in advanced disease without therapeutic alternative, previously received oncospecific treatment (surgical, radiotherapy, chemotherapy or the combination of these), ECOG ≤ 3, and normal functional parameters of clinical laboratory. The exclusion criteria was those patients who were receiving another investigational product or nimotuzumab, pregnant or breastfeeding, who refused to sign the informed consent and with ECOG > 3.

**Treatment**

Nimotuzumab was administered in a dose of 200 mg intravenously; once a week, for 12 weeks in the induction stage and 200 mg every 14 days until the onset of unmanageable toxicity or worsening of the general condition of the patient above 3 according to the ECOG scale. The safety and efficacy (concerning to survival time) associated with the treatment with the MAb was evaluated. Demographic and toxicity parameter were tabulated with descriptive statistics and was reported as absolute and relative frequency.

**Toxicity evaluation**

For the toxicity analysis, the adverse events were classified according to the table of Common Toxicity Criteria (CTC version 3.0 of the National Cancer Institute of USA). Tenor more times are considered as more frequently and as related to the AcM treatment all those had some causality relation with it (possible, probable and definite).

**Evaluation of effectiveness**

The main efficacy variable was the overall survival time (OS), measured from the date of initiation of treatment with the MAb to the date of death of the patient (or last life date news), using the Kaplan-Meier nonpara-

| Variables | Patients (%) |
|-----------|-------------|
| Age (years) | 65 (29-77) |
| Histological | |
| - Epidermal Carcinoma | 44 (58.6) |
| - Adenocarcinoma | 6 (8.0) |
| - Others | 25 (33.3) |
| ECOG | |
| 0 | 19 (25.3) |
| 1 | 37 (49.3) |
| 2 | 3 (4.0) |
| 3 | 5 (6.7) |
| Not available | 11 (14.7) |
| Previous Treatment | |
| CRT | 38 (50.7) |
| RT | 12 (16.0) |
| CT | 3 (4.0) |
| Not previous treatment | 12 (16.0) |
Results and Discussion

Seventy-five patients were included from January 2008 to May 2010, with follow-up until December 2018, the clinical, pathological and therapeutic characteristics are summarized in Table 1. The average age was 65 years (range of 29-77), the most frequent histological type diagnosed was squamous carcinoma in 58.6% of the patients and ECOG 1 predominated in 49.3%.

More than 50% of patients had locally advanced disease at diagnosis, however, at the inclusion in the clinical program were in advanced, persistence or relapse disease. Most of the patients received previous oncospecific treatment: 50.7% received the combination of external radiotherapy with Co60 in four pelvic fields for a total dose of 50.4 Gy with concurrent cisplatin chemotherapy at a weekly dose of 40 mg/m² of body surface for 5 weeks, followed by Brachytherapy of high dose rate with 600 cGy/session, four times. Twelve patients had not received previous oncospecific treatment because they were in very advanced clinical stages and refuse chemotherapy.

Toxicity

Table 2 shows the safety profile in the clinical study. Adverse events (AE) that appeared during the treatment in all of the included patients were evaluated as causal relationship. In 53 patients, 354 events were reported, out of 75 patients, treated 53 patients manifested some AE and only 18 were classified as related to nimotuzumab (24%), the most frequent reported SAEs were related to the administration of the MAb.

Although the pain was the most frequent reported associated with the administration of the MAb, this could be related to the advanced stage of the disease, since the intravenous administration require stay at the hospital for observation, at least two hours, however, this product was well tolerated and administered for prolonged periods in patients with long survival. Any new AE was reported for nimotuzumab [19-21]

The NCCN for cervical cancer in 2018 recommends as category 2B the use of Bevacizumab mAb as monotherapy [22] and, more recently, pembrolizumab [23,24], however, grade III adverse reactions such as hypertension, thromboembolic phenomena and gastrointestinal fistula are described [22]. Taking into account that the patients evaluated in this study received nimotuzumab as a second line therapy (or more) for their recurrent or metastatic disease and the administration was prolonged, even after progression in the monotherapy way, the safety profile is superior to other antibodies and are better tolerated.

In a prospective, phase II pilot study conducted from 2010 to 2015 in patients with locally advanced cervical carcinoma at Nanhai District University Hospital in the People's Republic of China, which included 80 patients treated with nimotuzumab concomitant with chemotherapy. Fourteen percent of the patients reported complete response, 88% control of the disease and 74% stable, showing a safety profile similar to that of the present study [17]. Another phase II study was reported by Cetina and cols. in 2015, conducted at the National Institute of Mexican Cancerology (INCAM) combining...

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### Table 2: Nimotuzumab toxicity. General profile of toxicity in the clinical study by patients and adverse events.

| Adverse Events (AE) | Nimo (%) |
|---------------------|----------|
| Patients with any adverse event | 53 (70.7) |
| Patients with any adverse event Related to the treatment | 18 (24) |
| Patients with any adverse event grade 3 and 4 | 34 (45.3) |
| Patients with any adverse event grade 3 and 4 Related to the treatment | 4 (5.3) |
| Patients with severe adverse Events | 14 (18.7) |
| Patients with any severe adverse Events Related to the treatment | 3 (4) |
| Adverse Events | 354 (100) |
| AE Related to the treatment | 83 (23.4) |
| AE grade 3 and 4 | 61 (17.2) |
| AE grade 3 and 4 Related to the treatment | 4 (1.1) |
| Severe Adverse Events (SAE) | 77 (20.1) |
| SAE Related to the treatment | 4 (1.1) |

SAE Patients (%)

| SAE Patients (%) | Total |
|------------------|-------|
| MILD | MODERATE | SEVERE | VERY SEVERE |
| PAIN TUMOR RELATED | 3 (7.0) | 9 (12.7) | 5 (7.0) | 2 (2.8) | 21 (29.5) |
| COUGH | 3 (4.2) | 4 (5.6) | 0 (0.0) | 0 (0.0) | 7 (9.9) |
| ANEMIA | 0 (0.0) | 3 (4.2) | 2 (2.8) | 0 (0.0) | 5 (7.0) |
| CYSTITIS | 3 (4.2) | 1 (1.4) | 1 (1.4) | 0 (0.0) | 5 (7.0) |
| ASTHENIA | 3 (4.2) | 0 (0.0) | 1 (1.4) | 0 (0.0) | 4 (5.6) |
| VOMITING | 1 (1.4) | 2 (2.8) | 0 (0.0) | 0 (0.0) | 3 (4.2) |
| OTHERS | 16 (22.5) | 10 (13.4) | 0 (0.0) | 0 (0.0) | 26 (36.6) |
| TOTAL | 31 (47.3) | 29 (40.8) | 9 (12.7) | 2 (2.8) | 71 (100) |

71 patients had some SAE related to nimo, considered more frequent when it appeared 3 or more times.

Of the 9 classified as severe, 2 were anemia, 1 asthenia, 5 pains, 1 cystitis and of the very serious 2 were pain.

There were only two reported rash with light intensity and no related deaths were report.
cisplatin/gemcitabine chemotherapy with nimotuzumab in 17 patients with persistent or recurrent cervical cancer after several lines of chemotherapy treatment. The induction phase of the treatment with the MAb for 4 weeks concurrent with the chemotherapy, and the maintenance every 14 days until tolerability of the patient. The treatment was well tolerated with mild or moderate toxicity, both in the induction and maintenance stages, achieving a stabilization of the disease in 35% of the patients included in the study [18]. Independently of the fact that in the present study the patients did not receive a concomitant oncpecific therapy such as those mentioned above, which could increase the toxicity of nimotuzumab, they were in a more advanced stage of the disease and even then the treatment was very well tolerated (Table 2).

**Efficacy evaluation**

Efficacy was evaluate in terms of overall survival, analyzed by intent to treat (ITT) and per protocol (PP) (Figure 1). Others factors that could have an impact on survival were also take into account for this analysis: first, the histological type (Figure 2); second, the ECOG at the time of inclusion (Figure 3). The behavior regarding previous therapies was consider as the third factor, resulting that untreated patients demonstrated a higher survival. This last could be related to the expression of EGFR in the untreated tumors, since the previously therapy could have a modification of the over expression on the tumor surface, as has been previously reported by several authors, [25-28] for this reason it is now essential to characterize the tumor once again when starting a biological therapy (Figure 4).

In the ITT survival analysis (OS) a median survival of inclusion was obtained at 23.5 month study, with a 4-year survival rate of 40.7%. It was higher than that reported in the GOG 240 study ( EC phase III random-ized study of Bevacizumab (anti-angiogenic MAb) con-comitant with chemotherapy with cisplatin/paclitaxel and topotecan/paclitaxel in which a significant OS of 17 months vs. 13.3 months was obtained (p = 0.004)) [29]. Those patients who received at least 6 doses of nimotuzumab (PP) did reach 31.1 months as median survival.

In the study conducted by Cetina and cols; where nimotuzumab treatment with concurrent chemotherapy with cisplatin and gemcitabine was applied to patients

![Figure 1: Global Survival of the study.](image_url)
with recurrent or persistent cervical cancer; the OS was 299 days (9.9 months) [18]. While in another similar investigation by Chen and cols. applied nimotuzumab with chemotherapy in similar doses and schemes at present, OS was obtained of 11.96 months (8.11-23.95) with an IC of 95% [17].

Recently Lu, et al. reported a clinical trial in 28 patients previous diagnosed of locally advanced cervical cancer. Neoadjuvant treatment of chemotherapy with concurrent platinum salts was applied with intensity modulated radiotherapy (IMRT) together with weekly nimotuzumab for 5-6 weeks was the treatment proposed; obtaining a complete response in 8 patients and a partial response in 20. It was possible to perform a hysterectomy with lymphadenectomy in 22 cases, demonstrating moderate over expression of EGFR moderately in 28.6% and high in 50% of cases [30].

In this combination modality of nimotuzumab with the onco-specific treatments of chemotherapy and radiotherapy before surgery, they reported a follow-up of 22 months with a range of 5 to 39 months, a survival of 90.0% [30].

The present report shows the results of nimotuzumab as monotherapy in patients with persistent, recurrent or metastatic cervical cancer, yielding a median OS of 27 months. The 48-month survival rate indicates that the probability of patients being alive at that time is upper than 40.0% for ITT and 50.0% for those who at least received the induction dose with nimotuzumab for 6 weeks. (Figure 1)

Some factors were taken into account to identify the populations that could benefit from this treatment. In this case, Figure 2 shows the histological type of the original tumor could be related to the advantages of OS, being the patients with epidermal carcinoma, the most benefited in relation to adenocarcinomas (ADC) and other tumors of epithelial origin that were included in the study. This could be directly relate to over expression of EGFR in squamous tumors due to its epithelial origin, much higher than ADCs of glandular origin [24-28,30].

In the evaluation of the general state of the patients at the moment of the inclusion, it influenced not only the extension of the disease, but also the sequels of the previously surgical procedures, chemo and radiotherapeutic treatments, which were reported as functional or organic, in different degrees before entering the study. Although all were included in advanced disease and without therapeutic alternatives, those that presented a better performance status (0-1) had a superior

| Histology          | Patients N=60 (%) | Mean (months) | Median (months) |
|--------------------|-------------------|---------------|-----------------|
| Epidermal Carcinoma (EC) | 44 (58,7%)        | 53,5 (39,6-67,4) | 27,0 (0,00-97,1) |
| Adenocarcinoma (ADC) | 6 (8,0%)          | 42,1 (4,6-79,6)  | 5,4 (0,00-27,4)  |
| Others             | 10 (13,3%)        | 30,6 (6,5-54,6)  | 13,8 (7,7-20,0)  |
| Global             | 60 (80,0%)        | 49,3 (37,3-61,2) | 20,9 (9,6-32,2)  |

**Figure 2:** Survival according to histological type at diagnosis.
Figure 3: Survival according to ECOG.

![ECOG Survival Graph](image)

| ECOG | N=46 | Mean (months) | Median (months) |
|------|------|---------------|-----------------|
| 0    | 19 (29.6%) | 62.8 (41.4-84) | - |
| 1    | 37 (57.8%)  | 80.43 (27.3-55.4) | 16.3 (1.9-3.8) |
| 2    | 2 (4.7%)    | 33.7 (0.0-71.7)  | 17.5 (0.0-40.4) |
| 3    | 5 (7.2%)    | 19.9 (0.0-53.9)  | 3.7 (0.0-8.9)   |
| Global | 46 (100%) | 46.8 (35.3-58.3) | 18.8 (8.8-28.8) |

P=0.013

Figure 4: Survival according to treatments prior to inclusion in the Expanded Use Program.

![Survival Graph](image)

| Previous Treatments | Mean (months) | Median (months) |
|---------------------|---------------|-----------------|
| No                  | 69.3 (53.1-85.5) | Not reach |
| Yes                 | 49.0 (37.4-60.7) | 23.2 (13.1-33.3) |
| Global              | 55.5 (44.6-66.5) | 27.0 (0.0-89.2) |

P=0.018
Dueñas and cols [34] Reported tumor resistance in - is developing every time that it receives an oncospecific could be related to the transformations that the tumor previous treatment showed a survival advantage, this the group is unbalanced, those that did not receive disease without therapeutic option. Independently that relation to those who arrived at the study in advanced count the patients who received previous therapies in 

Figure 3 shows the survival analysis taking into account the patients who received previous therapies in relation to those who arrived at the study in advanced disease without therapeutic option. Independently that the group is unbalanced, those that did not receive previous treatment showed a survival advantage, this could be related to the transformations that the tumor is developing every time that it receives an oncospecific therapy, in relation to the virgin tumors of treatment. Dueñas and cols [34] Reported tumor resistance increase as direct relation to the state of tumor progression because it being expected to acquire additional genetic and epigenetic alterations as it progresses, and this, directly or indirectly, provides protection from the cytotoxic or cytostatic effect of the onco-specific treatments used [33,34]. An additional factor that frequently interferes with the use of chemotherapy is the poor reserve of the bone marrow because of previous pelvic radiotherapy. This also worsened the response to treatments. Coincidentally, patients who did not receive previous treatments had tumors of the squamous cell carcinoma type, which are reported as EGFR over-expressers, so that therapy with an inhibitor should result in a therapeutic benefit [25,26].

Conclusions

The use of nimotuzumab as biological therapy in patients with cervical cancer in advanced stages proved to be safe and well tolerated. The survival profile obtained in the present study could constitute a therapeutic alternative in this stage of the disease whether persistent, recurrent or metastatic at diagnosis, in which no response has been obtained with the onco-specific treatments established for previous stages. The histological type, the previous treatments and the ECOG could be prognostic factors to the response to this therapy. It is proposed to continue a phase III clinical trial in the same set of patients or in earlier stages of this disease in which one use a nimotuzumab as a therapeutic approach.

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