Introduction

Tracheotomy and tracheostomy are medical devices used for many years. This technique was first described during the second century AD by Galien and Aretaeus of Cappadocia [1]. Even though a tracheostomy is performed only in an oncological purpose, there are many reasons of performing a tracheotomy (oncological or neurological fields, after a prolonged intubation in Intensive Care Units…). However those devices are still inspiring fear which is mainly caused by their symbolic impacts and the lack of training of healthcare professionals in use of such devices. After leaving their specialized center (surgery or rehabilitation center), patients and their caregivers are forced to fend for themselves or under care of inadequately trained medical and paramedical professionals. This means a lot of distress on the patient’s side but also on the health professional’s side and a frequent refusal of care.

Very little objective data concerning the care management of patients suffering from local habits are reported in the specialized literature.

AIM

The aim of this national working group was to formalize a consensus on care management (nursing techniques, management of complications) of patients with tracheotomy or tracheostomy at patients’ home and in care centers, whatever the cause of tracheotomy / tracheostomy is, benign or malignant.

Methods

Design

This study is a mixed research [2] combining both quantitative and qualitative researches. Furthermore, the methodology of a qualitative research with iterative focus groups was applied and the Consolidated Criteria for Reporting Qualitative Research (COREQ) checklist is reported in Table 1.

Setting / Participants

The group was supported by the French Association for Supportive Care in Oncology (AFSOS) and the Regional Oncology Network of Rhone Alps, Lyon, France. A national call for project was performed on Internet to recruit multi-professional volunteers at the beginning of 2015, anyone was accepted. Seven phone call with focus groups, each 2 hours, were set during 2015. Minutes performed during each meeting to modify the text

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**Abstract**

**Aim and objectives:** Formalize a consensus about patient’s care management (nursing techniques, management of complications) with tracheotomy or tracheostomy at home and in care center.

**Background:** Tracheotomy and tracheostomy are medical devices used for many years thus inspiring fear mainly by their symbolic impacts and the lack of training of health professionals.

**Design:** This study is a mixed research with a qualitative methodology including iterative focus groups.

**Methods:** A national call for project was performed on Internet to recruit multi-professional volunteers early 2015, anyone was accepted. Seven phone call with focus groups, each 2 hours, were set during 2015. Finally the project was presented and validated in “Guidelines and French Oncology Networks” J2R congress in Nantes, France in December 2015 and published on the web.

**Results:** Definitions, anatomic and physiologic notions, then the different types of cannulas are presented in the document. The management of tracheotomy/tracheostomy daily cares with protocols and videos, of the complications (infections, bleeding …), of functional sequelae (phonation, swallowing) at home and in care centers is then explained and demonstrated.

**Conclusions:** Finally, this work has led to a national consensus on the management, at home and in care centers, of tracheotomy and tracheostomy management and their potential complications.

**Relevance to clinical practice:** Such a work has never been done before. It aims to be comprehensive and didactic by means of figures and decision trees. This study will be useful and could be implemented despite local habits.
Table 1: COREQ checklist.

<table>
<thead>
<tr>
<th>Domain 1: Research team and reflexivity</th>
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<tbody>
<tr>
<td><strong>Personal Characteristics</strong></td>
<td></td>
</tr>
<tr>
<td>1. Interviewer/facilitator. Which author/s conducted the interview or focus group?</td>
<td>The coordination was performed by GB, HLC, LG</td>
</tr>
<tr>
<td>2. Credentials. What were the researcher’s credentials? E.g. PhD, MD</td>
<td>GB is MD, HLC is PharmD.</td>
</tr>
<tr>
<td>3. Occupation. What was their occupation at the time of the study?</td>
<td>GB is ENT surgeon in Valence, France. HLC and LG are project managers in the Regional Oncology Network of Rhône Alpes, Lyon, France</td>
</tr>
<tr>
<td>4. Gender. Was the researcher male or female?</td>
<td>GB is a male; HLC and LG are females.</td>
</tr>
<tr>
<td>5. Experience and training. What experience or training did the researcher have?</td>
<td>GB is an ENT surgeon specialized in head and neck cancers. HLC and LG are specialized in drafting care oncology guidelines.</td>
</tr>
<tr>
<td><strong>Relationship with participants</strong></td>
<td></td>
</tr>
<tr>
<td>6. Relationship established. Was a relationship established prior to study commencement?</td>
<td>A national call for project allowed recruitment of participants. Anyone was accepted. Except from GB and HLC, and HLC and LG, nobody knew each other at the beginning of the focus groups.</td>
</tr>
<tr>
<td>7. Participant knowledge of the interviewer. What did the participants know about the researcher? e.g. personal goals, reasons for doing the research</td>
<td>At the first focus group each participant was presented, nobody knowing each other. But the goal (formalize a consensus about care management) was known of each participant.</td>
</tr>
<tr>
<td>8. Interviewer characteristics. What characteristics were reported about the interviewer/facilitator? e.g. Bias, assumptions, reasons and interests in the research topic</td>
<td>During the initial presentation, GB indicated he knew the topic but not the methodology, contrary to HLC and LG</td>
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<th>Domain 2: study design</th>
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<tr>
<td><strong>Theoretical framework</strong></td>
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<tr>
<td>9. Methodological orientation and Theory. What methodological orientation was stated to underpin the study? e.g. grounded theory, discourse analysis, ethnography, phenomenology, content analysis</td>
<td>States of the art in the literature and in each participant’s health institution were performed</td>
</tr>
<tr>
<td>10. Participant selection</td>
<td></td>
</tr>
<tr>
<td>11. Method of approach. How were participants approached? e.g. face-to-face, telephone, mail, email</td>
<td>Seven telephonic focus groups took place during 2015. Notes after each focus group were integrated in the main document that was e-mailed before the next meeting for personal re-work.</td>
</tr>
<tr>
<td>12. Sample size. How many participants were in the study?</td>
<td>16 participants were recruited.</td>
</tr>
<tr>
<td>13. Non-participation. How many people refused to participate or dropped out? Reasons?</td>
<td>As they were volunteered, no participant refused to participate.</td>
</tr>
<tr>
<td><strong>Setting</strong></td>
<td></td>
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<tr>
<td>14. Setting of data collection. Where was the data collected? e.g. home, clinic, workplace</td>
<td>Data were collected in workplace</td>
</tr>
<tr>
<td>15. Presence of non-participants. Was anyone else present besides the participants and researchers?</td>
<td>Nobody was present besides the participants or researcher.</td>
</tr>
<tr>
<td>16. Description of sample. What are the important characteristics of the sample? e.g. demographic data, date</td>
<td>The main characteristics were a wide repartition around France and job diversity among participants.</td>
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</table>

**Data collection**

| 17. Interview guide. Were questions, prompts, guides provided by the authors? Was it pilot tested? | No interview guide existed. A general document for each care standard was done by the French Association for Supportive Care in Oncology (AFSOS) |
| 18. Repeat interviews. Were repeat interviews carried out? If yes, how many? | Seven telephonic conference calls took place in 2015 |
| 19. Audio/visual recording. Did the research use audio or visual recording to collect the data? | The research used no recording. |
| 20. Field notes. Were field notes made during and/or after the interview or focus group? | Notes were made during each conference call. |
| 21. Duration. What was the duration of the interviews or focus group? | Each focus groups lasted between 1.5 to 2 hours. |
| 22. Data saturation. Was data saturation discussed? | The data saturation was not discussed. |
which has been send by email to each participant before the next meeting.

Approval by every member of the focus group, another national call for the project was performed to recruit new readers to improve its quality. Then interregional guidelines have been reworked by new experts, submitted and approved during the “Guidelines and French Oncology Networks” congress in Nantes, France in December 2015. Finally they were published on AFSOS and regional oncology networks websites.

**Results**

**Theoretical considerations**

*Definitions, anatomic and physiologic notions: They are presented in Table 2.*

**Different types of cannulas:** Many types of cannulas exist. Eight different types can be described according to three options: with vs without (disposable or reusable) inner cannula (IC), cuffed vs cuffless and fenestrated vs non-fenestrated. For instance a cuffed fenestrated without (disposable or reusable) inner cannula (IC), cuffed vs cuffless and fenestrated vs non-fenestrated. For instance a cuffed fenestrated without (disposable or reusable) inner cannula (IC), cuffless vs fenestrated and non-fenestrated vs non-fenestrated. In case of fenestrated cannula (OC). It is therefore recommended to use IC for daily use.

Cannula without IC are common in intensive care units for patients who preferably be used when there is no risk of aspiration.

- Cannulas with and without IC

IC allows easy cleaning of sputum without removing the outer cannula (OC). It is therefore recommended to use IC for daily use.

Cannula without IC are common in intensive care units for sedated patients and when the caregivers are aware of the suction process and cleaning cares.

- Shapers, with a wide collar and without IC, are used after total (pharyngo) laryngectomy to avoid tracheostomy stenosis.

- **Cuffed and cuffless cannulas**

A cuff protects the lungs from fluid leakage. Therefore it is recommended if there is a risk of bleeding (just after the operation for instance), of swallowing disorder (after head and neck surgery or neurological diseases) or of major gastroesophageal reflux.

The cuff can be inflated temporarily in risk situations (swallowing for example) and deflated the rest of the day. When the cuff is inflated and unless the cannula is fenestrated a valve must not be absolutely used (risk of asphyxia).

The gold-standard method of cuff-inflation is a manometer control: the cuff must be 20 to 25-cmH2O air-inflated. Under than 20cmH2O, cuff is not sealed; over than 25cmH2O, i.e. more than capillary pressure, mucosal necrosis then stenosis are possible. When there is no manometer available (especially at the patient’s home), the necessary air volume of air is a necessary parameter which must be transmitted from the health institution nurse to the home nurse. Alternatively the cuff can be air-inflated while the patient is speaking. When the patient is not able to speak, the cuff is filled-in enough. Very anecdotally (hyperbaric chamber) the cuff can be inflated with sterile water (volume determined by the ENT surgeon). A special attention must be paid in case of accidental cuff rupture due to the risk of aspiration.

Cuffless cannulas are more lightweight, allow speaking and must preferably be used when there is no risk of aspiration.
- Fenestrated and non-fenestrated cannulas

A fenestrated cannula has a window top-directed, facilitating the path of the airflow through the larynx (Figure 1C). The main convenient cannula for that purpose is a cuffed fenestrated with IC system with patient who swallows the wrong way. Two IC are provided: a fenestrated one and a non-fenestrated one. When the patient wants to speak, even if the cuff is filled-in, the fenestrated IC must be used. When the patient wants to eat, the non-fenestrated IC avoids aspiration.

- Optional and peripheral systems:

  Fixation system: elastic, adjustable, comfortable but it present a risk of skin irritation, usually with Velcro® system. The fixation can also be facilitated by hooks. One man’s finger or two woman’s fingers should be put between the necklace and the neck. If this space is larger, the cannula may go out; if smaller it becomes too tight and uncomfortable.

  Chuck: systematically provided for cuffed and some cuffless cannulas, this device is used instead of the IC into the OC. It protrudes slightly and ends with a sphere. It enables the cannula set being very little traumatic and decreasing the risk of swallowing the wrong way outer of the trachea.

  Cone: fixed on the OC, it prevents the IC to move. The inner diameter is identical to the IC one, which allows to breathe easily.

  Half-moon device: fixed on the OC, it also prevents the IC to move. A deflector twice-reduces the diameter of inspired airflow but allows deflection of sputum.

Speaking valve: includes a membrane which is opened when the patient is breathing in and closes when the patient is breathing out. It allows speaking and the beginning of cannula weaning.

Filter: connected to the outer extremity of the cannula, it avoids dusts, humidifies and warms the inspired air. Optionally, it includes a metal or silicon structure which uses expired water vapor to humidify and warm the inspired air. If they do not prevent bacteria colonization, they do not increase its frequency or type [3].

Practical considerations

**Tracheotomy/tracheostomy daily care management:**

- **Cautions**

  Enteral feeding must be stopped two hours before the care and can begin just after.

  Individual protective equipment is mandatory.

  Information, reassurance of the patient and his closest entourage is strongly advised.
• Delays

In case of tracheotomy, from the operation to day +1 or 2, only the IC can be removed to be cleaned up, e.g. when skin is fixed around the tube. The neckline can also be prudently replaced if needed. At day + 2 or 3, the cuffed cannula can be replaced by a non-cuffed cannula (the presence of an ENT surgeon should be compulsory). A special training is then given to the patient and his entourage in order to become autonomous.

In case of tracheostomy, from the operation, the whole cannula can be removed to be cleaned up. The presence of an ENT surgeon is not mandatory but recommended. A special training is given to the patient and his entourage for autonomy as soon as possible.

Free videos are available for cuffed or uncuffed cannula removal.

(http://espacecancer.sante-ra.fr/Ressources/referentiels/videos/Changement%20de%20canule%20(2015)_720p.mp4) and cannula removal by an autonomous patient


• Cares

Local tracheotomy/tracheostomy care protocols must be done, describing every different cares. Habits are different from one country to another so a single protocol cannot be presented. Two French protocols are available for tracheotomy (http://espacecancer.sante-ra.fr/Ressources/Documents/Protocoles-soins/protocole_soins_tracheotomie_VF.pdf) and tracheostomy (http://espacecancer.sante-ra.fr/Ressources/Documents/Protocoles-soins/protocole_soins_tracheostomie_VF.pdf).

Infectious complications management: There is a continuum between colonization, tracheobronchitis and pneumonia and it is sometimes difficult to distinguish them.

The American Thorax Society [4], defines the as follows:

Colonization: isolation of a potential pathogen in tracheal microbiological cultures without purulent secretions or obvious clinical infection

Tracheobronchitis: clinical signs (fever greater than 38°C, purulent sputum) and biological signs (leukocytosis or leukopenia, positive culture of tracheal aspirates or sputum) without new or progressive radiographic infiltrate

Pneumonia: clinical and biological signs of tracheobronchitis and new or progressive radiographic infiltrate

• Types of microbiological cultures

The main problem here is the high rate of false positive because of tracheal colonization or contamination of the sample. Therefore bacteriological sampling techniques must be standardized [5,6]. For instance, endotracheal aspirate cultures give more micro-organisms than bronchoscopic ones.

Recommendations:

- The best sampling is bronchoscopic: bronchoalveolar lavage or protected specimen brush.

- Endotracheal aspiration is quite reliable when bronchoscopy is not available.

- Percannular sampling must not be performed.

- Quantitative cultures are compulsory. Semi-quantitative cultures are not reliable for diagnosis and qualitative cultures overdiagnose infections.

Positive thresholds of quantitative cultures are reported in Table 3.

• Tracheal colonization

Colonization is precocious (less than two weeks [7], even from the first day [8]), polymicrobial (aerobic, anaerobic [7,9,10] and fungal [8], in 80 to 100% of cases) [7,11,12].

The mechanisms of colonization are very close to those of chronic obstructive pulmonary disease or bronchial dilation [13]:

- Loss of natural protection by oral and nasal fence allowing direct contamination of microorganisms in the trachea and bronchi. There are very few correlations in microorganisms between the upper aerodigestive tract and the tracheobronchial tract [14-16].

- Decrease of mucociliary clearance causing the stagnation of sputum.

Most of reported bacteriology studies [7-12] are now outdated and microbial ecology has probably been improved since. Most frequent aerobic bacteria were Pseudomonas aeruginosa (in up to 75% of patients), Haemophilus influenzae, Klebsiella pneumoniae, Staphylococcus aureus, Serratia marcescens, Proteus sp, alpha-hemolysis Streptococcus group, Acinetobacter baumanii. Most frequent anaerobic bacteria were Fusobacterium nucleatum, Peptostreptococcus sp, and Bacteroides fragilis. Most frequent fungi were Candida (54.5%, half of which are C. albicans), Aspergillus fumigatus and Aspergillus flavus.

**Table 3: Positive thresholds of quantitative cultures. BAL = bronchoalveolar lavage, PSB = protected specimen brush, CFU = Colony-forming unit.**

<table>
<thead>
<tr>
<th></th>
<th>BAL</th>
<th>PSB</th>
<th>Endotracheal aspiration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Threshold</td>
<td>10³ or 10⁴ CFU/ml</td>
<td>10⁵ CFU/ml</td>
<td>Variable (in general 10⁵ CFU/ml)</td>
</tr>
<tr>
<td>Sensitivity</td>
<td>42 to 93%</td>
<td>33 to 100%</td>
<td>74 to 97%</td>
</tr>
<tr>
<td>Specificity</td>
<td>45 to 100%</td>
<td>50 to 100%</td>
<td>74 to 100%</td>
</tr>
</tbody>
</table>

**Commentaries**

- Beware of high tracheal tract contaminationsV (high level of squamous cell in direct examination). A direct examination can be done contrary to the two other samplings.

- Results are more specific than sensitive for pneumonia (i.e. a negative PSB is strong against the diagnosis of pneumonia). Beware of a high interindividual variability of sampling
• Recommendations:

A systematic bacteriological survey is strongly discouraged [9] because there is little agreement between the germs of colonization and those responsible for infections, or if it is the same seed, the antimicrobial susceptibility is often different, flora being unstable over long periods [13].

There is no interest in trying to eradicate the colonizing flora [4]. Indeed, after antibiotic treatment, microbiological samples are not sterilized with either the persistence of the same microorganisms or the appearance of new resistant germs [7,11,17].

• Tracheobronchitis and pneumonia

On average, the incidence of infectious episodes at home during one year (tracheobronchitis and pneumonia) was estimated at 10 per 100 person per year (83% of tracheobronchitis and 17% of pneumonia) [10].

There is no indication of probabilistic tracheobronchitis treatment because it is not an emergency. After bacteriological sampling, culture results must be awaited to prescribe a specific antibiotic therapy. Tracheobronchitis may improve spontaneously within 48 hours.

Pneumonia should be routinely treated with empiric antibiotic therapy according to environmental microbial ecology (intensive care unit vs general care unit vs home) secondarily adapted to the results of bacteriological samples.

Other complications management: They are summarized in Table 4.

Functional sequelae management: Phonation with a cannula

• Phonation with a cuffless cannula

Without dyspnea or breathing intolerance, a phonation valve can be adapted to the cannula (Figure 1A).

• Phonation with a cuffed cannula

Phonation is possible only in two limited circumstances:

- The cuff is deflated (Figure 1B):

Phonation is possible but more difficult than when the cannula is cuffless because the presence of the balloon, even deflated, reduces the free area between the cannula and the tracheal wall so the outgoing air flow.

- The cuff is inflated but the IC and OC are fenestrated (Figure 1C)

Apart from these two circumstances, a speaking valve must never be put on a cuffed cannula.

In order to avoid errors, most centers do not use phonation valve for cuffed cannulas.

Swallowing with a cannula: Swallowing troubles may be associated with the presence of the cannula because it prevents the rise of the larynx complicating swallowing. Some centers advise against swallowing with the presence of the cannula and prefer to wait for the cannula removal before swallowing. On the other hand, a swallow in the wrong way is obvious (externalization through the cannula) and the patient can be suctioned immediately.

<table>
<thead>
<tr>
<th>Table 4: Other complications management.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Problem</td>
</tr>
<tr>
<td>Difficulty in introducing the cannula within the trachea</td>
</tr>
<tr>
<td>Granuloma between IC and OC</td>
</tr>
<tr>
<td>Tracheal stenosis</td>
</tr>
<tr>
<td>Accidental removal of the cannula</td>
</tr>
<tr>
<td>Vomiting</td>
</tr>
<tr>
<td>Periorificial granuloma</td>
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<tr>
<td>Cannula obstruction</td>
</tr>
<tr>
<td>Pericannular bleeding</td>
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<tr>
<td></td>
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<td></td>
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<tr>
<td></td>
</tr>
<tr>
<td>Tracheal bleeding</td>
</tr>
</tbody>
</table>

Citation: Buiret G, Gautheron L, Labrosse-Canat H (2016) TracheotomY/Tracheostomy Management at home and in Care Centers. Arch Otolaryngol Rhinol Ear (2): 061-069. DOI: 10.17352/2455-1759.000027
Swallowing troubles can also be independent from the presence of the cannula. The causes are various and often linked to the indication of tracheotomy:

- Pharyngolaryngeal hyposensitivity abolishing swallowing reflexes (prolonged intubation)
- Swallowing desynchronization
- Laryngeal disclosure
- Pharyngeal propulsion default (sequelae of surgery or radiotherapy)
- Delay in the opening of the upper esophageal sphincter

An ENT examination including a neurological examination and a fiberoptic laryngoscopy (with food testing and subglottic fiberoptic control by the tracheostomy hole) is a valuable aid for speech therapy which is the base of rehabilitation (relearning swallowing praxis, postures, food textures...).

- Monitoring of patients with tracheotomy / tracheostomy: Respiratory function:
  - Respiratory amplitude, heart rate, agitation, sweating, pulling signs, SpO2, free airway
  - Cuff pressure, better with a manometer. The cuff sealing must be daily verified: the inspired volume in the cuff must be equal to that which was injected. Otherwise, the balloon is surely become porous and the cannula is to change.
  - Nature of tracheal aspirations: quantity and appearance of secretions (search for possible infection)

Fixing the cannula and tolerance of this fixation.

Home coming: Home coming must be anticipated in order to identify the patient needs and to arrange education of the patient and his entourage.

- Multi-daily cannula cares must be organized with suctions, changes and/or cleaning of the cannula, aerosols...

Material Management must be delivered and tested before the patient is returning home:

- Two tracheal vacuum suctionsing systems including one with battery
- Equipment maintenance, cleaning, storage and verification
- One emergency cannula and one with smaller diameter in reserve
- Other equipment if needed (oxygen, aerosol, respirator)

Pharmacists or home health providers are essential to care.

Procedures to follow in case of problems must absolutely be known by the patient, his entourage and his caregivers: bleeding (Figure 2), accidental decannulation (Figure 3), and cannula obstruction (Figure 4).

Discussion

The routine management of patients with tracheotomy or tracheostomy is problematic because of the lack of training of the patient, his entourage and the caregivers. The different devices provided with a cannula and their functions are sometimes not even known by the caregivers. After a reminder of the technical considerations, the different materials available and their indications,
the article exposes a progressively obtained multi-professional and pragmatic consensus about care management of patients with tracheotomy or tracheostomy.

The design of the study was first discussed:

- A review of literature was an incomplete method because very little or no reference is available.
- A conventional quantitative research was not indicated. Indeed there was no data and the study design wasn’t subject to statistical assumptions and conditions.
- The aim of a qualitative study is to understand and interpret social interactions. The main purpose is the study of human and cultural impacts on a particular subject. However the data format of a qualitative study with iterative focus groups and field notes was interesting.
- A mixed research appeared to be particularly adapted. From an epistemological point of view, the purpose of our work corresponds to a pragmatic justification (what works for whom in specific contexts) [2]. Another interest was the flexibility in the study design: participant responses affected how and which questions researchers asked afterwards. A mixed research also allows connection between theory and practice.

As consequent we decided to use a mixed research for the purpose of the study using rigorously the methodology of a qualitative research (reported in Table 1). The national validation of the consensus followed two additional steps after the focus groups:
A national diffusion for proofreading, the collection of data and consideration of remarks
- The reviewing with a group of experts during the "Guidelines and French Oncology Networks" congress.

The principal limitation of the present study is that very few references are available in the literature. Another limitation is the date of the publication especially bacteriological ones. By cons care have not changed since the routine use of tracheotomy.

Conclusion

After a schematic presentation of medical devices (types, functions, criteria of choices...), this work has led to a national consensus concerning the management at home and in care centers, of tracheotomy and tracheostomy and their potential complications.

Relevance for Clinical Practice

Such a work has never been done before. It aims to be, comprehensive and didactic with its figures and decision trees. This study will be useful to decline in each country in accordance to local habits.

Acknowledgements

The authors specially acknowledge Hubert Riccardi from the Réseau Régional de Cancérologie Rhône Alpes and Céline Labrosse for the English translation and all the contributors of the focus group of the Association Francophone des Soins Oncologiques de Support.

References


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