

Mapping out the social experience of cancer patients with facial disfigurement

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ABSTRACT

This article contributes to the limited literature on the social consequences of cancer generated facial disfigurement by reporting the result of an exploratory analysis of interaction between facially disfigured cancer patients and strangers and acquaintances (secondary groups). Secondary groups are those in which membership occurs due to performance of formal and/or non-intimate roles. Interaction is studied as it takes place in different social settings. Individuals who are affected by cancer of the head and neck region can now expect to survive for many years after the cancer is detected and later surgically removed. Because of surgery, these survivors live the rest of their lives with facial disfigurement and are stigmatized and socially excluded. It follows that a new and socially relevant situation has emerged: as medicine develops and allows more patients to survive, it forces them to spend significant portions of their lives dealing with the stigma associated with facial disfigurement. Research on social issues pertaining to facially disfigured cancer patients remains sparse. Limited knowledge has been produced on the “social context” within which interaction between the disfigured and relevant social groups takes place. To date most research has focused on the individual and his/her ability to adapt to the condition of facially disfigured. To address this scientific gap and document the manner through which the interaction process is socially created and evolves, interviews with fourteen facially disfigured cancer patients were carried out. These interviews were designed to reconstruct the interaction experiences of these individuals in different social contexts. Data were analyzed through the qualitative approach of grounded theory. Results indicate that patients can be divided into two groups: *Occasionally Comfortable Patients* and *Always Comfortable Patients*.

Occasionally comfortable patients are individuals who experience different levels of comfort in interaction. In some situations they do not feel stigmatized, but other interactions constitute the contexts within which this discomfort emerges. Discomfort in interaction was employed as an indicator of stigmatization. Interacting groups were divided into small and large. Intrusion (unsolicited attention to patients) in interaction in large and small groups always generates uncomfortable situations. Sympathy (unsolicited comments and/or actions in support of patients) is associated with comfort in interaction in small groups and produces varying patterns in the case of large groups. Benign neglect (a situation in which interacting individuals do not pay particular attention to patients) produces comfort in interaction within large groups and varying outcomes in the case of small groups. Always comfortable patients are those who do not experience discomfort in interaction regardless of the size and characteristics of the interacting group. The article concludes by stressing that facially disfigured cancer patients should be prepared to face different interaction patterns. Simultaneously, efforts should be made to educate patients and the general public about these interaction patterns.

Keywords: Facial Disfigurement; Cancer; Stigma; Social Interaction

1. INTRODUCTION

The objective of this article is to illustrate the results of an exploratory analysis on the patterns of social interaction experienced by individuals who are facially disfigured because of cancer. Individuals who are affected by cancer can expect to be cured and/or survive for a number of years after the cancer is detected and later surgically removed [1,2]. This is also the case of patients with head and neck cancer who remained facially disfigured

because of it [3,4]. In this particular instance, surgical intervention signifies the removal of portions of the face that are affected by the malignancy. Additional consequences are possible and they often include the collapse of parts of the face that may not be directly touched by surgical intervention [5]. Accordingly, patients' facial disfigurement may involve a larger portion of the face than that originally affected by cancer [6]. Surgical procedures to correct these alterations are common along with the availability of increasingly sophisticated prostheses [3]. Yet, results often do not rectify notable differences from the "normal" face. Accordingly survivors typically live the rest of their lives with major alterations of their normal facial appearance.

In contemporary society, the face represents one of the most notable items employed to determine identity and participate in social interaction [7-9]. It is a central element of communication [10,11], an item employed to attribute both "normality" and ownership of socially desirable characteristics [12-16] and a feature that defines interaction as individuals endowed with a pleasant face are better treated than other and less attractive members of society [17,18]. Owing to the social importance of the face, facially disfigured individuals are stigmatized and experience difficulties when interacting with other segments of society [11,12,14-16,19-21]. It follows that a new and socially relevant situation has emerged. As medicine develops and allows more patients to survive, it also makes them spend significant portions of their lives dealing with the stigma associated with facial disfigurement.

Research on social issues pertaining to facially disfigured cancer patients remains sparse and attracts even less attention than the already limited research associated with other forms of acquired and congenital disfigurement [10,15,22-24]. Additionally, analytical problems have limited the accuracy of available research results [25]. In this context, limited knowledge has been produced on the "social context" within which interaction between the disfigured and relevant social groups takes place [21,25]. Through this paper we would like to contribute to this limited literature. In particular, this article offers the results on an exploratory analysis on patterns of interaction between disfigured patients and members of secondary groups (acquaintances and strangers) as interaction occurs in different social settings. Interaction with other groups, such as family members and friends (primary groups), is important. However, in this work we focus exclusively on interaction between patients and members of secondary groups. The study of the interaction with other groups remains a relevant topic to be further explored.

Methodologically, the study consists of in-depth interviews with fourteen cancer patients who underwent surgery to treat head and neck cancer and remained facially disfigured because of it. Patients were enrolled following

a protocol that excluded minors and those who could not express themselves in English. Given the relative limited size of this patient population, the adoption of a qualitative approach was recommended. A semi-structured questionnaire was employed and data were analyzed employing *Grounded Theory* [26]. Grounded theory is a technique of qualitative data analysis that allows the identification of patterns directly from the data. Rather than hypothesis verification (pre-existing ideas to be verified in reality), grounded theory allows the development of categories (variables) and their relationships (patterns) that are not established a priori. These categories and patterns emerge directly from the data and can be always traced back to the data themselves. Because there are no pre-conceived hypotheses to be tested, this technique is particularly appropriate for exploratory analyses [26]. This is a qualitative study as it is aimed at illustrating the manner through which the patterns of interaction unfold. It explains the "how" these patterns manifest themselves and develop rather than how often they occur in reality. It also stresses the process through which the interaction is created and grows rather than its outcomes. Given the small size of the sample, this approach has been considered more appropriate than quantitative statistical analysis.

The statements concerning the result have been verified through established data verification techniques. In particular the techniques of "saturation" and "negative cases" were employed. In the case of saturation, the creation of categories is verified by searching for possible instances in which data cannot be explained by the category in question. When no more instances to be explained by the category in question are found in the data, the category is said to be saturated and therefore verified. In the instance of negative cases, categories are compared with situations in which their attributes are contradicted by evidence. As categories that have negative cases are eliminated, the remaining ones are considered verified. These techniques were applied to the construction of the categories and patterns employed in this article.

The objective of the article is to identify patterns of interaction between facially disfigured individuals and members of secondary groups. Secondary groups were defined as non-intimate, formal groups consisting of acquaintance and strangers. Results indicate that facial disfigured patients feel stigmatized when interacting within small and large groups and the interacting counterpart(s) shows "intrusion" (*i.e.*, unrequested attention to patients including unwanted questions, remarks and stares). Patients respond in varying manner when "sympathy" (*i.e.*, unsolicited comments and/or actions showing support) and "benign neglect" (*i.e.*, no particular attention paid to patients) characterized the interaction. A group of patients that do not feel stigmatized during various types of interaction was identified. The paper

concludes by stressing the importance of studying stigmatization through a sociological approach and by pointing out that cancer survivors and the general public alike could benefit from education about these interaction outcomes.

2. CANCER AND FACIAL DISFIGUREMENT: SALIENT LITERATURE

Stigma is a mark of disgrace attached to people who are considered different. As indicated by Goffman [13], difference is socially constructed and is the outcome of discrepancies between an individual virtual social identity (expectations about what that individual ought to be) and his/her actual social identity (the attributes he/she actually possesses) [13]. When the actual social identity is perceived as departing from normality, the individual is “reduced in our minds from a whole and usual person to a tainted, discounted one. “Such an attribute is a stigma” [13]. Stigma is attached to an individual’s feature “that is deeply discrediting” and that separates that person from the group of the normals. However, its actual genesis should not be linked to attributes but, rather, to the interaction between the stigmatized person and other members of society. “A language of relationships, not attributes, is really needed” Goffman states [13]. Stigma is generated by the existence of a number of blemishes. There are those of individual character such as homosexuality, dishonesty, imprisonment, radical political behavior, and addiction. There are those of tribal stigma that are related to a person’s religion, ethnicity or race. And there are those of “abominations of the body” that refer to physical abnormalities. Goffman includes facial disfigurement in this category [13].

Following the seminal work of Erving Goffman [13], social stigma has been widely studied and this production includes works such as those on stigma generated from diseases (*i.e.*, cancer and AIDS) [27], physical disabilities [28,29], and mental health¹ [30,31].

In spite of this wealth of contributions, the stigma caused by facial disfigurement has been the subject of only a relatively small number of works [10,15,21,22, 24,32]. These analyses stressed the social importance of the face and the problems that affect those who display visible facial blemishes [12-16]. In a society in which

¹This abundant literature has also underscored important limits of the use of the concept of stigma. For instance, stigma has been studied with a strong individualistic focus, it is often employed by people who do not belong to stigmatized groups, there is no consensus on a common definition, and the existence of these multiple definitions allow the charge that this concept is too inclusive to be actually informative ([28] Cahill and Eggleston, 1995:682; [34] Link and Phelan, 2001: 365-366). There are also uncertainties about its manifestations as “felt” stigma—the individual’s “shame” associated with the blemish—is much more common than the rare “enacted” stigma—or the existence of overt episodes of discrimination ([35] Jacobi, 1994:269).

individuals are fully clothed for virtually all of their social activities, the face represents one of the most notable physical attributes and a significant source of social information prior to, and during, social interaction [8,9, 30,33]. Accordingly, people possessing an attractive face² are not only considered physically pleasing³, but they are often viewed as endowed with intellectual and emotional characteristics that are unrelated to their physical appearance [10,11,36]. Intelligence, kindness, likableness, and high morality are frequently associated with an attractive appearance. Unsurprisingly, these individuals are also better treated by others than less attractive members of society [17,18]. Lacking some of these physical attributes, facially disfigured individuals commonly engender negative responses by other members of society. Stigmatized and socially excluded, their ability to interact is often distorted and interaction is the source of problems including verbal and physical abuse, ridicule, hostile behavior, and isolation [10,12,37].

The literature identifies two types of facial disfigurement: congenital and acquired. Acquired disfigurement received less scientific attention than the already limited literature on congenital disfigurement [24]. Acquired disfigurement is further divided into trauma- and cancer-generated disfigurement. Research demonstrates that individuals with acquired facial disfigurement suffer psycho-social consequences that are different and, at least to some degree, more pronounced than those experienced by individuals with congenital disfigurement [18,24]. Among individuals with acquired facial disfigurement, cancer patients⁴ experience less social and psychological problems than individuals who have been disfigured because of trauma [38]. However, cancer patients’ “fear of dying is immense” [39]. And this situation affects these individuals and their family members’ perception of disfigurement [40]. In this context, patients are more preoccupied with the evolution of their cancer than with the social consequences of the scars that the disease left on their faces [24,40]. As this fear of dying diminishes, however, the process of dealing with the deformity affects both patients and family members

²Early research on physical appearance paid more attention to the face than the rest of the body. It was stressed that the face was the most visible part of the body, its stability of appearance was greater than body appearance both in terms of the individual’s lifespan and developmental stages, and physical attractiveness was predominantly determined by the face. Recent research, however, while still emphasizing the face [4], has underscored the growing importance of the body in the determination of beauty and attractiveness [9,20].

³*e.g.*, “Beauty is perceived as residing principally in the face.” [33].

⁴While the social perception of cancer has changed in recent decades, this disease engenders a wide variety of attitudes and responses that differentiate it from other pathological situations [46]. Often, these attitudes and responses are stigmatizing [47,48]. However, differences have been recorded between reactions to forms of cancer that are perceived as “uncontrollable”—such as breast cancer—and those that are perceived as “controllable”—such as lung cancer due to smoking. Because the latter are seen as deriving from the patient’s voluntary actions, more stigmatizing reactions are expected [49].

[39,40]. The association of cancer and disfigurement is persistent. In effect, therapy almost inevitably mandates surgical removal of cancer-affected parts of the face making it an undesirable consequence of successful medical intervention [20,41,42].

Among the limits of this otherwise important literature is the lack of attention paid to the social context within which the consequences of facially disfigurement emerge [12,21,25,32]. Emphasis is placed on the individual, his/her characteristics, and efforts to adapt to his/her condition of disfigured [4,39]. This posture downplays “the everyday experience of the disfigured population in social settings” [17,21,25,32,37,43]. In this context, the “interaction process” through which disfigurement is experienced is viewed almost exclusively in terms of the significance of individual behavior as a predictor of social outcomes [19,32,43]. This is also the case when other social actors—such as partners—are studied [37,44]. In particular, limited attention is paid to the fact that disfigurement is socially constructed and generated through processes of interaction that involve multiple actors and take different forms according to the settings in which they unfold [21].

The individual remains the research focus also in studies on the relationship between cancer-generated facial disfigurement and stigmatization [45]. Indeed, research on stigma has been “decidedly individualistic in focus” [34]. Accordingly, the ways in which interaction between the facially disfigured and the “normals” unfolds in different realms of society has been understudied. As Thompson and Kent put it “Most studies have examined the ‘view from the inside,’ with little work [carried out] on the social and the ‘view from the outside’” [25].

In this literature, the interaction of disfigured individuals within primary groups has been privileged over than within secondary groups [15,16,45]. In particular, the spheres of the “family” and “caregivers” are portrayed as social settings in which the disfigured find more comfortable terms of interaction. The family is seen both as a “safer place” in which the disfigured feel protected and supported and the institution that provides them with alternative messages than the stigmatizing ones coming from society and its strangers [15,16]. Interaction with caregivers, and in particular those between surgeon, social workers and disfigured patients, is also viewed as comfortable [15]. To be sure, race-, ethnicity-, and culture-based forms of discrimination in the hospital are documented [16] and improvement in the manner in which health care professionals deal with those who are facially disfigured is frequently identified as a primary programmatic objective. Nevertheless, the hospital is often identified as one of the settings in which patients achieve comfort in interaction [15].

The same literature identifies secondary groups in

terms of “strangers” who populate “public places” where they not only intrude into the lives of the facially deformed through “staring, remarks, and questions, or obvious eye avoidance [16], but also negate that “civil inattention” normally granted to other members of society [16]. These groups constitute the “popular mind” [16] that produces “generalized prejudice” [16]. The social settings where secondary groups exist are seen as generating stigmatization as people deal “differently with those who have undergone facial surgery” [15] and engender “fear” to the disfigured [15]. Ultimately, society is the source of generalized stigmatization through widespread stereotyping that allows the “...unsightly face [to be] utilized as a visible symbol or a personification of evil, disease, criminality, or mental deficiency” [16]. This process of stigmatization is so pervasive that it is viewed as if it is conducted uniformly throughout society regardless of actors and settings [15,16]. While evidence indicates that society is the primary source of stigmatization [10,20,24,37,39], the manner through which facially deformed cancer patients experience stigmatization in different spheres of society is understudied [45]. van Doorne and his associates [39], share the view that members of the immediate family provide strong support to cancer patients who are facially disfigured. They also stress that strangers are consistently the source of stigmatization. In their view, further investigation is needed to ascertain the manner in which non-immediate family members and acquaintances react to facially disfigured patients.

In essence, the study of stigmatization of the facially disfigured is couched in approaches that stress the actions and adaptability of the individual, on the one hand, and the homogenous dimension through which stigmatization is created in society, on the other. In this context, the manner in which facially disfigured cancer patients experience stigmatization while interacting with secondary groups in different spheres of society has not been adequately explored. This literature does not differentiate between the many settings that form the sphere of society and that are the contexts in which interaction with secondary groups takes place [45]. Owing this gap in the literature, the objective of this paper is to provide an exploratory analysis of the patterns of social interaction experienced by individuals who are facially disfigured because of cancer as they interact with members of secondary groups.

3. METHODS

This exploratory research employed in-depth interviews with fourteen individuals who were facially disfigured because of head and neck cancer. A purposive sample was selected from a list of patients who were registered at MD Anderson Cancer Center in Houston, Texas. This

particular group of patients had cancer of the upper face area and specifically orbital and periorbital cancer. Potential participants were identified following the recruitment guidelines. They were adult patients (18 years old and older), who were diagnosed with various forms of head and neck cancer, underwent facial surgery to remove the malignancy and, in the process, acquired observable changes in their facial appearance. Additionally, they were required to be able to communicate in a meaningful manner with investigators.

In-depth individual interviews were conducted by phone between January 2008 and April 2009. Twenty potential participants were contacted. Three of them could not participate for scheduling problems and three refused to be part of the project. Finally, a total of eight men and six women were interviewed. The median age of these patients was 66 years and the youngest patient was 31 and the oldest 81. At the time of the interview, the post-surgical period ranged from ten months to thirty five years and the median was five years. Some patients underwent additional reconstructive and plastic surgical procedures. While the extent of disfigurement varied, all of the patients were left with significant alterations in their facial appearance.

A semi-structured questionnaire was employed. Questions covered topics such as the individual’s demographic and socio-economic characteristics; experience with cancer and treatment; perceived attitudes and actions of others; and interaction in various events and places. Patients were interviewed by a single research team member. This procedure was adopted to assure consistency in the interview procedure. With the patient’s consent, interviews were audio recorded and written notes were also taken.

The transcribed texts were analyzed through grounded theory methodology [26,50]. This methodology permits the identification of conceptual categories and patterns among them from the text. Categories and patterns were verified through the techniques of “saturation” and “negative cases.” Secondary groups were defined as nonintimate, formal groups formed by strangers and acquaintances. Following data analysis, salient interaction patterns were identified and patients were divided into those feeling comfortable, uncomfortable or having varied or mixed feelings. The feeling of uncomfortable was considered an indicator of the patient being stigmatized either in terms of felt stigmatization or enacted stigmatization. Felt stigmatization refers to the perception of the patient. Enacted stigmatization refers to overt actions of stigmatization. Interaction patterns were identified and divided into three categories. *Intrusion* indicates that people pay unsolicited attention to patients, ask unwanted questions, make unwelcome remarks and stare. It also indicates that they make their unspoken curiosity felt. *Sympathy* refers to unsolicited comments and/or

actions showing support to patients and the desire to be of assistance. *Benign neglect* denotes a situation in which interacting individuals did not pay particular attention to patients. Size of interacting group was divided into small and large. These concepts indicate that the secondary group in which interaction took place was either large or small. The actual size of the group was determined through patients’ own descriptions of the settings.

4. FINDINGS

Following grounded theory analysis of the data, two groups of patients were identified: *Occasionally Comfortable Patients* (N = 10) and *Always Comfortable Patients* (N = 4). Patterns of interaction with *Occasionally Comfortable Patients* are visually synthesized in **Figure 1** and basic descriptive statistics for this group are presented in **Table 1**. Basic descriptive statistics for *Always Comfortable Patients* are summarized in **Table 2**.

Over 71 percent of the patients studies suffer from some forms of stigmatization. In relative terms, men suffer from stigmatization slightly more than women (75% versus 66.7%). Yet, women over 55 years of age experience stigmatization in slightly higher terms than men of the same age group. Intrusion is the source of stigmatization for the majority of patients and impacts men and women equally. Older men experience stigmatization from sympathy more than women of the same

		Size of Group	
		Small	Large
Types of Interaction Postures	Intrusion	Uncomfortable	Uncomfortable
	Sympathy	Comfortable	Comfortable Uncomfortable
	Benign Neglect	Comfortable Uncomfortable	Comfortable

Figure 1. Patient-secondary group interaction by size of group

and types of responses to disfigurement for Occasionally Comfortable Patients.

Table 1. Basic characteristics and patterns of interaction of Occasionally Comfortable Patients, N = 10 (total patients studied N = 14).

Gender	Men 6 75%	Women 4 66.7%	Total 10 71.4%
Age and Gender	Men > 55 4 66.7%	Women > 55 3 75%	Total 7 70%
	Men < 55 2 33.3%	Women < 55 1 25%	Total 3 30%
Intrusion	3 50%	2 50%	
	Men > 55 2 66.7%	Women > 55 2 100%	5 50%
	Men < 55 1 33.3%	Women < 55 0 0%	
Sympathy	2 33.3%	1 25%	
	Men > 55 1 50%	Women > 55 0 0%	3 30%
	Men < 55 1 50%	Women < 55 1 100%	
Benign Neglect	1 16.7%	1 25%	
	Men > 55 1 100%	Women > 55 1 100%	2 20%
	Men < 55 0 0%	Women < 55 0 0%	

Table 2. Basic characteristics of Always Comfortable Patients, N = 4 (total patients studied N = 14).

Gender	Men 2 25%	Women 2 33.3%	Total 4 28.6%
Age and Gender	Men > 55 2 100%	Women > 55 2 100%	Total 4 100%

age group while older women experience benign neglect relatively more than men. Due to the limited size of the sample and the qualitative design, these statistics are reported only as an illustration of possible quantitative trends. More pertinent findings are presented below in the discussion of the qualitative analysis.

4.1. Occasionally Comfortable Patients

Occasionally comfortable patients are individuals who

experience different levels of comfort in interaction. In some situations they do not feel stigmatized, but other interactions constitute the contexts within which this discomfort emerges. Intrusion in interaction in large and small groups always generates uncomfortable outcomes. Here patients are stigmatized. Sympathy is associated with comfort in interaction in small groups and produces varying patterns in the case of large groups. Benign neglect produces comfort in interaction within large groups and varying patterns in the case of small groups.

4.1.1. Intrusion

This is a situation in which patients feel uncomfortable interacting within small and large groups alike. Members of interacting groups grant disfigured individuals the particular status of “different” and proceed to construct an interaction in which this condition is reinforced. Patients are noted because their different appearance. They are drawn in interaction patterns in which their status of different is constructed and reinforced through unwanted attention. People’s questions, stares, remarks constitute elements that build a view of disfigurement that is stigmatizing. While varying, patients’ responses are characterized by dissatisfaction with the interaction. Interaction in large and small groups alike provides the terrain for the development of this constructed stigmatization.

4.1.1.1. Small Groups

Joy provides an account of this type of interaction with small groups. She says:

“Sometimes I notice that people...can tell that this side of my face does not make the same movements [than the other side]. They are probably aware that there is something strange about that woman...The first time that I took my six year old to dancing—my six year old granddaughter, she dances every Wednesday and then she is at church afterwards. The children have special things going on after dancing at the church... My daughter works a lot and so that Wednesday is mine. I take them and do all of that kind of thing. Well, the first few times when I was taking her to dancing, I sit and wait... can’t go off and leave her, wait in the room with the other parents and I could tell that some of the mothers were looking at me strangely, not in a mean way or anything like that but more curious, like there’s something strange about that woman. What could it be? And generally, I have found that if I feel like somebody is making me feel uncomfortable... I’ll go ahead and say something.”

She adds:

“I went to a wedding shower for one of my daughter’s friends who I didn’t know very well and there were a lot of young women there that I did not know well... and I was uncomfortable there. I felt like the majority of them probably did not know that I had this problem [cancer generated facial disfigurement] and I’m sure they were wondering... that was an uncomfortable day.... I didn’t

really enjoy that.”

4.1.1.2. Large Groups

Arthur provides an instance of the interaction between patients and large secondary group. He says:

“I feel like everybody’s always staring at me. It is strangers. [I have negative feelings] when dealing with strangers. Probably the worse is in restaurants. Because you are sitting there for a long time and the people around you are all eventually looking at you or you feel they are. [It is also the same] when you walk past somebody in a store or on the street or something... They stare at you... It is very crude and insensitive and it makes me skittish towards other people... it makes me feel very uncomfortable...I don’t go out very much anymore.”

Janet adds:

“When people keep staring at me and pretend they are not looking at me... It bothers me. Most adults—they will look at me. They all notice, I’m sure. A lot of them try to pretend... But it happens everywhere. The worst is at places like Super Centers. That where I found that people are most rude. It also bothers me that people let their kids be so rude, particularly big kids.... They run around you in front of me and go ‘she doesn’t have and eye! Look, look!’ Then I want to smack them. I want to tell them ‘you are very, very rude.’”

4.1.2. Sympathy

Sympathy creates interaction patterns that are dissimilar for small and large groups. In small groups, data reveal that sympathy engenders comfortable interaction between disfigured patients and acquaintances. The latter’s statements and actions are intended and perceived as supportive and contribute to the construction of a situation in which disfigured patients find themselves at ease. The analysis indicates that varying patterns define the interaction between patient and members of large groups. Sympathy shapes positive interaction as people’s support is employed to construct advantageous conditions for patients. Instrumentally, patients use this support even in situations in which it is not needed. Simultaneously, patients also feel uncomfortable when sympathy transcends the necessary. They feel that their disfigurement is constructed as a grater disadvantage than it actually is and/or a situation that creates undeserved respect. In this instance, overt sympathy becomes a negative factor in the construction of interaction. It is important to note that sympathy is also associated with the existence of cancer. In some instances, interacting individuals display sympathy to patient primarily because of their status of survivors.

4.1.2.1. Small Group

Lisa provides evidence of a positive interaction with small groups. She says:

“Nobody tells me anything. The only thing is they ask me what happened, you know, what is wrong. That’s the question. Then I have to tell them that I had surgery and I had cancer and I had a tumor... This is friends, friends that I haven’t seen in a long time and I happen to meet in a store while I’m shopping, people that I haven’t seen in a long time... they will ask questions... and when I explain to them what happened everybody is nice. They just want to know what had happened to me and they don’t make [comments or remarks], not at all, everybody is nice.”

4.1.2.2. Large Groups

Sympathy in interaction with large groups engenders a variety of results that include both comfortable and uncomfortable situations.

Tom states: “Depends on whether or not I’m wearing my prosthesis, the black patch or the medical patch and I get different reactions for each of those circumstances. When I’m wearing my prosthesis out in public... they do not know that I had surgery. If I wear it ... on the street, in the store, in front of a group of people, at church or in a meeting, if they do not know I had surgery they still do not know I had surgery. [Because it is difficult to wear my prosthesis continuously]⁵, I wear my black patch. When I wear my black patch, I get better service. If I ride a plane... people treat me differently... they are sympathetic, overtly so... This is a person with a visual problem, we’ll treat him differently and they do.... I think they feel sorry for me... They seem to be more sympathetic towards me and more ready to help me in stores, primarily in store settings or at the library or other public places. When I’m out in public service areas, the service that I receive appears to be somewhat more magnanimous... I’m going to take advantage of this thing as much as I can.”

But he also indicates: “Sympathy is good, but at times I feel that I don’t need it at all. I can do things on my own and do not need any help.”

Mary says: “I’m very self-confident ... around others because more and more people are very receptive of cancer survivors. So if you had cancer and you are a survivor, then people are very supportive.... I could be a horrible person (I’m not a horrible person) and they would be like ‘oh, that’s ok if you are a cancer survivor.... People at work have been very supportive... nobody has been negative about my appearance.... They simply ask what happened to you.”

4.1.3. Benign Neglect

⁵This patient informs us that wearing the prosthesis creates a number of problems. There are problems associated with skin irritation where the prosthesis attaches to the face. When the skin is irritated, the discomfort forces the patient to wear either a black patch or a medical patch. Additionally, there are problems associated with the patients’ activities. Wearing the prosthesis causes severe headaches when traveling by plane. Discomfort is also felt when pursuing outdoor activities or exercising.

Benign neglect refers to situation in which interacting individuals did not pay particular attention to patients. In the case of interaction with small groups, data show that varying patterns of interaction are possible. In some instances, lack of interests on the part of interacting individuals is felt and acted upon in positive terms. Disfigured patients do not feel stigmatized. Simultaneously, though, similar situations engender opposite interaction patterns as patients feel uncomfortable in the absence of attention paid to them. The fact that interacting individuals “ignore” patients is not necessary an indication of comfort in interaction. In the case of interaction with large groups, benign neglect generates comfortable interaction patterns. The size of the group is a condition that allows patients to pass unnoticed and this situation facilitates the manner through they interact with this group of strangers.

4.1.3.1. Small Groups

Following are a few instances of the consequences of benign neglect in interaction with small groups.

Carl says: “I have not noticed anything. I don’t think people feel sorry for me any more or any less. I think that still they see me as Carl... When I’m with a small group of people, I don’t see anyone doing anything differently for me... They treat me the same as always... When I go shopping people look at me... it doesn’t feel good but I don’t let it bother me, I don’t feel like I’m less of a human. I don’t have to look down like I’m ashamed to look them in the face, or like they are going to reject me or anything like that.... It bothers me though.”

Ron states: “I generally try to avoid being seen in a prominent position... I really do believe that I have learned to accept it [my disfigurement], but I do realize that it’s noticeable to people and that it’s pretty obvious. ... I felt good about myself before my cancer. Now, I look in the mirror and I’m missing a part that’s very important to your looks...people at work tell me ‘I don’t even notice that your eye is missing, because I’m so used to you.... It bothers me, but I accept it.”

4.1.3.2. Large Groups

Joy provides an instance of benign neglect in interaction with large groups. She says:

“We [my husband and I] go to [college] basketball games. We have a lot of those here. At the basketball game is different because everyone is so intent on what [the players] are doing what they are watching that they do not pay attention to anybody, because I mean, they are glued on what is going on on the court so I don’t notice people reacting to me.... I’m pretty confident interacting with people except when I need to meet new people... I used to love meeting new people. I don’t really enjoy that so much anymore... but I like to be in a large crowd when people don’t notice you.”

4.2. Always Comfortable Patients

Always comfortable patients are individuals who do not experience problems interacting in small and large groups alike. They are well adjusted to their disfigured status and, while they encounter behavior that could lead to enacted stigmatization, the interaction does not affected by them. As episodes of enacted stigmatization are quire rare, the lack of felt stigmatization denotes non-stigmatizing interaction patterns. Basic statistics about this group are summarized in **Table 2**. As it can be seen from that table, no relevant differences seem to exist in terms of patients’ age and gender.

Fred says: “I feel no difference the way that people treat me... I don’t see any difference... when we go to the shopping mall, when we go to the restaurant, when we go anywhere. I see no difference in the way I’m treated now and the way I was treated five years ago [before surgery].... I have no problems with the way I look whenever I’m meeting anyone. It’s not something that I think about and it’s something that I don’t even consider.... I get some sympathy because I’m a cancer survivor, but this has very little to do with the way I look.”

Frank continues: [when I interact with people I’m very confident] Oh, yeah, I think so. I went back to work on a full-time job where I used to travel a lot and go to a contractor’s facility in [name of city]... all the time and meet with people, have meetings, give briefings and so, ... I don’t have any issue with it and there never seemed to be an issue with those people I deal with.”

[When I go out] “the only people that tend to react are kids. And they’ll look at you, and it is just like at my wife’s church preschool class. I went to see her at her class today and went to lunch after that, and the kids came up and just looked and said, hey, how you doin’? And she said, oh, my husband’s a pirate... they said, okay, great!... I haven’t really had a negative situation so far.... When I had my knee operation the anesthesiologist that did the knee operation came to my house during Halloween and I answered the door and he was standing there with his child, and he said: you’re really getting into the part, aren’t you? Thinking that I had an eye patch on for Halloween. I said, nope, I really have to have an eye patch. He said: Oh, I’m sorry. That’s the guy I had four months later to be my anesthesiologist on my knee surgery, and he mentioned how he had met someone with an eye patch. I said that was me. He said: Oh, I feel so bad about it. I said: don’t worry, it happens all the time. You know, you get dressed up for something or whatever... but... it doesn’t bother me and in fact I called the guy about a part on a pool the other day, told him my name, he said: yeah, you are the guy with the bad eye – wait, you don’t mind that I said that, do you? And I said, no, I don’t mind that you said that.”

5. CONCLUSIONS

In terms of the interaction between facially disfigured cancer survivors and secondary groups, this study identified the existence of two groups of patients. When interacting with strangers and acquaintances these cancer survivors can be grouped into *always comfortable patients* and *occasionally comfortable patients*. Always comfortable patients are patients who do not experience problems because of their disfigurement and their interaction with small and large groups does not lead to felt stigmatization. While episodes of enacted stigma may occur, they tend not to affect these well adjusted individuals, their counterparts, and interaction.

The group of patients classified as occasionally comfortable presents more complex patterns of interaction with secondary groups. This group experiences uncomfortable and stigmatizing interaction patterns in specific instances. At the same time, they experience non consistent or comfortable (*i.e.*, not stigmatizing) patterns of interaction in other situations. This is the group of patients that requires the most attention as their interaction with secondary groups varies according to the size of the group.

These findings add to the existing literature in a number of ways. First, they indicate that strangers do display varying patterns of interaction with the facially disfigured. Accordingly, the statement that “the reaction of people on the streets or in the neighborhood is ... consistent: they stare at most patient with facial defect” [39] (see also [15,16]) needs to be revised. This study demonstrates that the behavior of strangers is much more complex than previously recorded and it entails a number of distinct interaction patterns.

Second, this research clarifies questions raised by previous research on the interaction between facially disfigured cancer survivors and acquaintances [39]. Data on the various patterns of interaction within small groups demonstrate the complexity of interaction patterns existing between patients and acquaintances. Ultimately, acquaintances are the source of stigmatization and support alike as they intrude in the everyday existence of patients but also provide sympathy to cancer survivors.

Third, these findings contribute to the understanding the issue of the relationship between disfigurement and cancer. The existence of cancer mitigates the negative effects of facial disfigurement. It creates a process of interaction in which people display support to patients because they suffered from cancer. Once people learn that the patient’s facial disfigurement is cancer related, they tend to be overtly supportive regardless of other conditions.

Finally, this research underscores the importance of the reading of the social consequences of cancer generated facial disfigurement in relational terms. The social

consequences of facial disfigurement emerge as outcomes of interaction processes. As such, they involve patients and other segments of society as they interact in various social spheres. While the study of the manner through which individuals respond to disfigurement remains important, it is through the process of interaction that stigmatization or lack of it are constructed.

Three overall recommendations emerge from the analysis presented in this study. First, patients should be warned about the occurrence of possible difficulties when interacting in large and small groups alike. In this respect, they should be prepared to the possibility of experiencing episodes of intrusion, sympathy and benign neglect when interacting with small and large groups of acquaintances and strangers. Second, it appears desirable that interacting groups be sensitized about the patterns of interaction identified above. While complex, this solution should be considered as educational programs are discussed and implemented at various levels. Finally, the issue of the social consequences of cancer generated facial disfigurement should be studied with an interdisciplinary approach. In this context, it would be desirable to include a sociological component that would allow investigators to explore the collective dimension of the social construction of facial disfigurement and stigma.

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