

See discussions, stats, and author profiles for this publication at: <https://www.researchgate.net/publication/269037283>

Towards a classification of alexithymia: Primary, secondary and organic

Article · March 2014

CITATIONS

90

READS

3,499

3 authors:



Antonino Messina

18 PUBLICATIONS 164 CITATIONS

SEE PROFILE



Janelle N Beadle

University of Nebraska at Omaha

39 PUBLICATIONS 923 CITATIONS

SEE PROFILE



Sergio Paradiso

Institute of Cognitive and Translational Neuroscience (INCyT), INECO Foundation, ...

106 PUBLICATIONS 6,774 CITATIONS

SEE PROFILE

Towards a classification of alexithymia: primary, secondary and organic

Verso una classificazione dell'alessitimia in primaria, secondaria e organica

A. Messina¹, J.N. Beadle², S. Paradiso^{3,4}

¹ General Medicine Trainers, Health Department of Sicily, Catania, Italy; ² University of Iowa, Iowa City, USA; ³ Universidad "Diego Portales", Santiago, Chile and ⁴ University of Hawaii at Manoa, Honolulu, USA

Summary

Background

Emotion processing is essential for well-being and psychosocial adaptation. Alexithymia is widely viewed as an impairment in emotion processing that includes difficulty identifying and describing emotions. While there is a significant understanding of primary alexithymia, which is thought to be the result of developmental genetic and familial factors, secondary and organic alexithymia are beginning to be focus of research.

Method and results

The present review of the literature suggests the importance of differentiating between primary and secondary alexithymia, and the organic subtype of secondary alexithymia. Secondary alexithymia is thought to be a consequence of psychological stress, chronic disease, or organic processes (e.g. brain trauma or stroke) that occur after childhood (whereas primary alexithymia

is a developmental phenomenon). Organic insults to the brain may bring about the organic form of alexithymia by altering cerebral structures involved in emotional processing (e.g. anterior cingulate cortex, frontostriatal networks, callosus corpus, right hemisphere cortex and amygdala). The usefulness of differentiating among alexithymia categories and their relationships with brain structures known to subserve emotional processing is discussed.

Conclusion

We propose that differentiating between primary, secondary and organic alexithymia may potentially serve to develop better treatments for alexithymia.

Key words

Alexithymia • Classification • Brain injury • Emotion

Introduction: origin of the concept of alexithymia

Alexithymia is a psychological construct broadly describing individuals with deficits in emotion processing and awareness¹. Individuals who score high on measures of alexithymia show difficulty distinguishing emotions from bodily sensations, discriminating between cognition and emotions, and describing and communicating emotions to others². Harvard psychiatrist Peter Sifneos first used the term alexithymia (a = lack; lexis = word; thymos = emotion) to describe individuals who appeared, "different, alien beings, having come from an entirely different world, living in the midst of a society which is dominated by feelings" (Goleman³, p. 51).

Classically, alexithymia has been defined to include multiple facets including⁴:

1. difficulty identifying and distinguishing emotions from bodily sensations;
2. difficulty describing and verbalizing emotions;

3. poverty of fantasy life;
4. externally oriented thinking style;
5. poor empathizing.

During dynamic psychotherapy, the capacity to symbolize and translate emotions into language is often considered to be critical for improving symptoms. These capacities are greatly reduced or lacking in alexithymia. Sifneos' interest in identifying individuals with alexithymia was motivated by the goal to select individuals who would show improvement with short-term psychodynamic therapy⁵. The negative effect of alexithymia on psychotherapy has been empirically demonstrated in group, psychoanalytic and supportive psychotherapies⁶. The reaction of the therapist to patients with alexithymia may also have a negative effect on treatment⁷.

Dating back to earlier research, the body or "soma" was found to play a critical role in alexithymia⁸. The present review will describe the two-way relationship between alexithymia and the body, beginning with a discussion of historical views purporting that emotions are embodied

Correspondence

Sergio Paradiso, Neuroscience and MR Research, Department of Medicine, John A. Burns School of Medicine, University of Hawai'i, 1356 Lusitana Street, University Tower, 7th Floor, Honolulu, HI 96813, USA • E-mail: sergiop@hawaii.edu or paradiso.sp@gmail.com

experiences. The potential utility of distinguishing between primary, secondary and organic alexithymia based upon aetiology and proposed brain mechanisms will be described.

Some historical views on emotion and the body

Ancient Greeks and Romans traditionally viewed emotions as “perturbationes animi” (literally, disturbances of the soul), or modifications of the mental state of an individual not guided by reason (Cicero 1 sec. BC)⁹. Today, in light of progress in cognitive and affective neuroscience, the dichotomy between emotion and reason is less tenable¹⁰. The ability to perceive, modulate and express emotions are core cognitive features of emotional *intelligence*¹¹. Knowing one’s own emotions is an essential ability that promotes adaptive decision-making and goal-directed behaviour¹². Emotional awareness has been considered to be a prerequisite for managing bodily drives and delaying gratification, while contributing to the selection of emotionally adaptive behavioural responses¹². The philosopher and biologist Herbert Spencer (1820-1903) posited that an emotion is a subjective mental state of pain or pleasure associated with bodily manifestations (“the inner and the outer face of the same change” Spencer, 1855, p. 128)¹³. Spencer believed that in the animal kingdom emotions play a role in evolution and adaptation, permitting a comprehensive classification of impressions and inducing adaptive responses to specific situations¹³. Darwin’s theory of emotions¹⁴ expanded upon Spencer’s ideas and emphasized the concept that an emotion is an automatic response with stereotyped bodily changes. Darwin believed that an emotion was a mental state causing somatic effects and observed similarities between somatic expressions of emotions among humans and other animals (e.g. gnashing teeth in anger)¹⁴. Darwin distinguished basic emotions including joy, shame, anger, and disgust from social emotions such as love or hate that generally are less stereotyped and more complex¹⁴.

William James and the Danish physician James Lange viewed emotions as a link between the perception of an event and the consequent behaviour: emotion (from Latin *e-moveo* = to move towards) compels to act¹⁵. According to James, an emotion is the consequence of an activation of the neurovegetative system¹⁵. The perception of an

aroused neurovegetative system generates emotion (Fig. 1). To expand, it is James’ opinion that the behaviour that follows a perception is the emotion. He wrote: “the bodily changes follow directly the perception of the exciting fact, [...] our feeling of the same changes as they occur is the emotion”^{15 16}.

In contrast with James, Cannon viewed bodily expressions and cognitive appraisal of emotions as parallel processes¹⁷. This signifies that Cannon believed that the neurovegetative changes were not the cause of the emotion. Cannon highlighted that the perception of an emotional event (e.g. danger) produces activation of subcortical brain structures thus leading to a generalized sympathetic response. Activation of cortical structures is necessary for the conscious representation of the stimulus and its emotional tagging¹⁷. The late Magda B. Arnold (1903-2002) (American psychologist based at Loyola University, Chicago) thought that emotions are produced during a dialectical process occurring between the mind and external objects. She thought that emotions are products of our evaluation of events (the “appraisal theory of emotions”). In addition she showed that (usually), positive emotions generate approach behaviour while negative emotions generate withdrawal. Hence, emotions are not exclusively inner processes, but are generated by interactions between the subject (with her body) and the object¹⁸. Conscious representation of an event permits an evaluation of the basic perceptual data and may allow modification of the emotional response¹⁸. In this model, Arnold posits the pivotal role of emotional awareness in controlling emotion responses and emotion-activated behaviour. In spite of the critical differences between various theories of emotion, this very brief account of the history of emotion theory demonstrates the importance of the emotion-body connection.

Social role of emotion and alexithymia

Amidst the complexity of human societies and groups, emotions play an important social role. Basic emotions including fear, for instance, often promote affiliation among individuals, as frightened individuals may seek support from the group and utilize these group resources against a real or imaginary enemy¹⁹. Other emotions (for instance empathy or envy) are eminently social in nature¹⁹. Decreased levels of empathy have been linked to

EVENT > NEUROVEGETATIVE AROUSAL > EMOTION

FIGURE 1.

James-Lange’s theory of emotion. *La teoria delle emozioni di James-Lange*.

greater loneliness²⁰. Because of the importance of emotions both for the individual and social groups, the fact that some individuals show a diminished capacity to recognize and describe emotions has been widely regarded as a highly important psychological and clinical issue²¹. Individuals with moderate to high scores of alexithymia show an array of difficulties in their relationships with others, including interpersonal ambivalence, need for social approval and poor sociability^{22,23}. Interpersonal difficulties can cause emotional suffering and may prompt individuals with alexithymia to seek psychotherapy. However, individuals with alexithymia may not receive the full benefit from dynamic, supportive, or group, psychotherapy because poorer access and reporting of personal emotions are negative prognostic factors in psychotherapeutic treatment⁶.

Alexithymia: cognition, somatic symptoms and disease

Alexithymia and cognition

Individuals with alexithymia typically have difficulties using language to describe their experiences of emotions that are rooted in bodily sensations. This view is consistent with empirical studies showing the association between alexithymia and language difficulties²⁴. For instance, Henry et al.²⁵ observed a significant inverse correlation between difficulty identifying emotions and verbal fluency in patients with a history of traumatic brain injury. Furthermore, Onor et al.²⁶ showed that alexithymia was associated with weaker functioning (albeit not in the impaired range) in several cognitive domains including language, attention, memory, visual spatial abilities, and working memory. In a different sample of healthy adults, the Paradiso lab showed that the severity of alexithymia was associated with relatively poorer cognition, in particular in the domain of executive function²⁷. Thus, the extant literature suggests that one mechanism for alexithymia is cognitive with relative language and abstract reasoning and symbolization deficits²⁸.

Yet, the mechanisms of alexithymia may be more complex and may extend beyond traditional cognitive functions. As evidenced by research linking alexithymia to awareness of personal emotional and brain regions supporting it (see below)²⁹, poor emotional self-awareness and poor abstract thinking may underlie a diminished capacity to symbolically express emotions or alexithymia³⁰.

Alexithymia, disease, and somatic symptoms

The relationship between alexithymia and reported or medically ascertained physical symptoms is complex. Rather than having no emotional life, individuals with alexithymia communicate their emotions using somatic

channels³¹. In addition, individuals with alexithymia show an excess of medically ascertained physical illness³² and alexithymia among individuals with ascertained medical conditions may hinder recovery and delay rehabilitation efforts³³. Understanding the complex relationship between poor emotion processing and physical symptoms includes understanding that individuals with alexithymia tend to complain of body ailments and disturbances independently from the actual presence of somatic illness³¹. This issue is complicated by evidence of individual differences among people with alexithymia. Bermond³⁴ separates alexithymia into two types. Type I alexithymia shows affective and cognitive alterations. People with Type I alexithymia display poor awareness and expression of emotions. On the other hand, Type II alexithymia shows normal emotion awareness, but poor emotion expression, and these individuals are more prone to somatization than Type I³⁵.

Alexithymia is also associated with visceral hypersensitivity. There is evidence relating impaired processing of emotions with functional gastrointestinal disorders³⁶. This finding is consistent with the notion that physical symptoms in alexithymia may be related to somatosensory amplification (SA). SA has been defined as: i) excessive attention and hyper-vigilance to somatic symptoms; ii) exaggerated sensitiveness to physical sensations; iii) misinterpretation of physical sensations interpreted as a sign of disease³⁷. Somatosensory amplification³⁸ and somatization^{31,39} are generally associated with difficulty identifying and describing feelings (but less so with externally oriented thinking style).

However, not all research supports the association between alexithymia and somatic complaints. Some authors believe that alexithymia and somatization are two different conditions that are only sometimes associated⁴⁰. For example, children in particular show multiple somatic complaints without difficulty in describing or identifying feelings⁴¹.

Alexithymia and psychopathology

Before addressing the differences between primary and secondary alexithymia, a brief account on the association with psychopathology is needed. Alexithymia is a risk factor for suicide in subjects with brain injury⁴². This finding is an important reminder of the vast literature concerning the relationship between alexithymia and depression. This issue has been widely debated in the field. Briefly, researchers now believe that a positive association between alexithymia and depression exists (i.e. the greater the alexithymia scores, the greater the depression scores). Longitudinal studies have also uncovered persistence of some degree of alexithymia following remission of depression⁴³. Thus, alexithymia has both state-depen-

dent (e.g. mood but also general psychopathology²²) and trait-dependent features⁴⁴. Trait dependent features, association with personality traits (including avoidant, schizotypal, dependent and passive-aggressive)^{22 45} and lack of association with histrionic features⁴⁵ reinforce the notion that the construct of alexithymia captures psychological dimensions related to individual differences. Alexithymia is also found in other psychiatric conditions including autism, eating disorders and schizophrenia, all of which show degrees of deficits in social cognition^{20 46 47}. Based on these data, it comes as no surprise that alexithymia together with personality disorders⁴⁸ is a predictor of poor outcome in depression⁴⁹.

In the end, some have suggested that it may be useful to distinguish between depression with strong and mild alexithymia features⁵⁰, but further study in this area is needed. If strong alexithymia prevents a consistent report of emotion changes such that patients admit to sad mood only erratically, depression may take the phenomenological form of depression without sadness (or non-dysphoric depression)⁵¹⁻⁵³.

Primary versus secondary alexithymia

The need for distinguishing between primary and secondary alexithymia emerged rather early in the literature⁵⁴, but it took some time before this distinction was widely recognized in the field. Most studies that have contributed to the complex body of knowledge on alexithymia have examined individuals with primary alexithymia. Alexithymia is considered to be primary when emerging “as a life-long dispositional factor that can lead to psychosomatic illness” (Lesser⁵⁵, p. 533). Primary alexithymia may derive from a psychic trauma occurring during childhood⁵⁶ or from negative primary caregivers interactions⁵⁷. It has been recently suggested that genetic polymorphism of the 5-HT transporter-linked promoter region (i.e. L/L alleles) may influence the occurrence of alexithymia⁵⁸. Hence, primary alexithymia is currently thought of as a more or less stable personality trait that becomes molded during childhood and early adult years. Therefore, primary alexithymia is developmental in nature. It also has no purported organic or psychological risk factors (excluding those occurring in childhood, see discussion below).

Secondary alexithymia is posited to arise not during development, but as a consequence of events occurring later in life. These may be events with psychological significance and/or medical-surgical events (illnesses or disease)⁵⁹ that have a direct or indirect effect on brain functioning. Therefore, secondary alexithymia may have both psychological and/or somatic (organic) mechanisms⁶⁰. As evidenced briefly above, in addition to somatic symptoms, alexithymia can also be associated with mental

illness. Thus, whereas *primary* alexithymia may play a role as a vulnerability factor for mental illness, *secondary* alexithymia is thought to be a consequence of the illness⁶⁰.

If the stressful event is an illness (one that has no obvious direct consequences on brain functioning such as a hip fracture), alexithymia has been seen as a defense mechanism in an attempt to cope with the stress of the medical illness⁵⁴. Based upon the study of alexithymia in 53 inpatients examined in a teaching hospital psychiatric consultation service, researchers suggested that “alexithymia in the medically ill may play a defensive role as a state reaction” (Wise et al.⁵⁹, p. 287). In other words, alexithymia secondary to a psychologically significant event may be construed as a defense or protection against highly emotional events. This view is supported by the higher levels of alexithymia found in holocaust survivors⁶¹ and sexual assault victims⁶².

In summary, while primary alexithymia is widely thought to be a personality trait, in which affective processing is less developed than normal due to childhood trauma or genetic predisposition (e.g. polymorphism of 5-HT transporter-linked promoter region), secondary alexithymia is a condition occurring later in life either due to psychological trauma, or as a direct insult to brain regions supporting emotion processing and awareness⁶³. Note that this proposed sharp dichotomy is useful to frame the field. Clinical experience illustrates that determination of the primary or secondary nature of alexithymia may be in some cases debated. Examples may be youth with severe illness (alexithymia can be seen as secondary because of the illness or primary due to its ensuing during development) or alexithymia resulting from prolonged emotional stress during early development (e.g. hyperprotective parental bonding and specifically excessive maternal protection) mediating development of personality disorders⁶⁴. Therefore, primary and secondary alexithymia may be better viewed as extremes of a continuum, while individual decisions on aetiology are better left to the experienced clinician.

Organic alexithymia

As studies began to examine the hypothesis that alexithymia may be associated with localized brain damage⁶⁵, the observation that alexithymia may occur “de novo,” as a consequence of brain injury generated extensive interest in the field⁶³. The term organic alexithymia refers to a condition in which alexithymia is purportedly caused by organic damage to brain structures involved in emotional processing through indirect or direct insults to the brain. We suggest that it is conceptually useful to categorize organic alexithymia under the rubric of secondary alexithymia, with the understanding that further outcome

TABLE I.

Neuroanatomical substrates of primary, secondary and organic alexithymia. *Substrati neuroanatomici dell'alestitimia primaria, secondaria e organica.*

Brain region	Primary alexithymia	Secondary alexithymia	Organic alexithymia
Anterior cingulate cortex	Borsci et al., 2009 ⁷⁸ (decreased grey matter volume); Heinzel et al., 2012 ¹²⁷ (functional impairment)	Messina et al., 2011 ⁷² (decreased oxygen tension) (purported as for the location)	Sturm et al., 2011 ⁹⁰ ; Paradiso et al., 2008 ⁸⁹ (decreased grey matter volume)
Corpus callosum	Romei et al., 2008 ¹⁰⁸ ; Tabibnia and Zaidel 2005 ¹¹⁶ ; Lumley, et al., 2000 ¹¹⁴ ; Parker et al., 1999 ¹²⁸ (interhemispheric transfer impairment)		TenHouten et al., 1986 ¹²⁶ (cerebral commissurotomy)
Basal ganglia	Lee et al., 2011 ¹⁰¹ (functional impairment in caudate and frontostriatal circuitry)		Huang et al., 2012 ⁶⁶ (damage of basal ganglia after carbon monoxide poisoning)
Right temporal lobe	Borsci et al., 2009 ⁷⁸ (reduced grey matter in right temporal lobe)		
Right hemisphere	Lumley et al., 2000 ¹¹⁴ (functional impairment)		Spalletta et al., 2001 ⁶⁵ (right hemisphere stroke)

and treatment studies may indicate that this should be considered altogether as a separate category. Organic alexithymia is often more resistant to treatment⁶⁶, is not associated with a specific pre-morbid personality and is associated with cognitive impairment.

As briefly mentioned above, alexithymia observed among patients with medical illness represents a clinical puzzle. The question refers to the extent to which in a given patient alexithymia is a primary personality feature or a phenomenon secondary to the stress of the illness or a direct or indirect brain alteration (induced by the medical condition). Determination of the nature of alexithymia may be inferred from the clinical history (e.g. asking a next of kin on personality features prior to illness or injury), and from characteristics inherent to the alexithymia syndrome (e.g. cognitive impairment).

Studies have examined the possibility that alexithymia stems from direct brain damage. Spalletta et al.⁶⁵ observed that alexithymic characteristics were associated with right hemisphere damage among patients with stroke consistently with the role of right hemisphere in emotion processing^{67,68}. Becerra et al.⁶³ reported on a 21 year old man who developed alexithymia after a motor vehicle accident that caused damage to the orbitofrontal cortex, a brain region playing a role in decoding emotional significance of an event and strongly connected with many limbic regions including the amygdala^{10,69}. Subsequently, a study examining 54 individuals (67% men) with brain injury (which had occurred about 30 years prior) confirmed the strong association between brain injury and alexithymia (odd ratio = 2.64)⁷⁰. The

majority (61%) of subjects with history of traumatic brain injury early in life developed alexithymia, which in turn is a risk factor for suicide in subjects with history of traumatic brain injury⁴². The literature also shows that hypoxic lesions of bilateral globus pallidus provoked by carbon monoxide poisoning were related associated with a severe and resistant form of alexithymia⁶⁶. As reported by the Paradiso lab, a 44-year-old man showed impairment in emotional awareness after anoxic lesion of globus pallidus⁷¹. Another study that may be relevant for this discussion showed, low levels of haemoglobin in oncologic patients were directly associated with higher level of alexithymia⁷². The authors speculated that low oxygen pressure level may modify the functioning of the anterior cingulate cortex, a region critical for emotional awareness and particularly susceptible to perfusion changes⁷².

Whereas the existing literature on treatment of secondary and organic alexithymia remains wanting, it may offer guidance in some cases. For instance, improvement in alexithymia was found when 64 psychiatric patients underwent group therapy⁷³. Other studies show efficacy of the serotonin-norepinephrine reuptake inhibitor (SNRI) venlafaxine (75-150 mg daily) on alexithymia in patients who suffered a stroke⁷⁴.

In summary, there may be at least two types of secondary alexithymia. Specifically, one type develops as a consequence of psychosocial stressors and the psychological (e.g. defence) mechanisms set forth to attempt to cope with the stress⁷⁵, while the other (organic) occurs due to a direct or indirect insult to the brain. As such, organic alexithymia may follow alterations to brain regions sub-

TABLE II.Alexithymia: classification based on clinical history. *Alessitimia: classificazione in base alla storia clinica.*

	Primary alexithymia	Secondary alexithymia	Organic alexithymia
Aetiology	Genetically mediated or familial	Subsequent to psychological distress	Subsequent to trauma, vascular or other brain damage
Onset mechanisms	Early onset	Early or late onset	Early or late onset
	Sociocultural education, socio-economic status	Linked to chronic disease (e.g. cancer, diabetes, Parkinson's disease)	Linked to brain damage in regions subserving emotion processing
Treatment response*	Psychotherapy treatment resistance	Somewhat responsive to psychotherapy or pharmacological treatment	Responsive to pharmacological treatment
Outcome*	Poorer	Better	Better

* Based on research studies and personal experience of one of the authors (SP)

serving emotional awareness or participating in linking stimuli from the body to their abstract (language) conceptualization including neurodegenerative disorders (e.g. frontal temporal dementia, semantic dementia, Alzheimer's disease, cortical basal degeneration/progressive supranuclear palsy).

Neuroanatomy of alexithymia

Multiple brain structures have been posited to be involved in the pathogenesis of alexithymia (Table I). There is a broad overlap of brain structures purportedly involved in alexithymia independently whether this may be conceived as primary or secondary (including organic). What appears to be the case is that the involvement of one or another brain region does not allow distinction between primary and secondary or organic alexithymia. The distinction continues to rely on clinical assessment (Table II). Regions purportedly associated with alexithymia have included the anterior cingulate cortex, frontal striatal networks and basal ganglia, insula and amygdala, corpus callosum.

Anterior cingulate cortex

Since Papez's initial postulation (1937), consensus has broadened on the role of anterior cingulate cortex (ACC) in emotion perception and regulation⁷⁶⁻⁷⁸. Briefly, the ACC shows at least two functional parts: a rostral region playing a role in emotion processing and interconnected with multiple limbic regions; and a dorsal region connected with the lateral prefrontal, parietal and supplementary motor cortex showing greater specialization for attention and executive functions⁷⁹⁻⁸⁰. Rostral and dorsal ACC regions are closely connected allowing integration between the experience and the mental representation

of an emotion⁸¹. "Anatomical and functional continuum rather than segregated operations" between cognition and emotional components of the mind may take place in the ACC (Messina et al.⁸², p. 1). The importance of ACC in alexithymia may be attributed to the presence of spindle-shaped Von Economo's neurons. These may represent a *trait d'union* between perception and emotions⁸³. Von Economo neurons appeared in late phases of evolution when they took a role in the regulation of social and emotional functioning⁸⁴. The presence of these neurons is related to absolute brain size⁸⁵. Homo sapiens has a larger brain size (about 1500 cc on average) than homo Neanderthal (1400 cc on average)⁸⁶.

Von Economo's neurons (VENs) are bipolar neurons allocated in layer V of ACC and frontoinsula cortex in humans⁸³. These neurons were also found in animals with high social structure as apes, dolphins⁸⁷, elephants⁸⁵ macaques monkeys⁸⁸. The importance of Von Economo's neurons in social awareness, empathy and self-referential processing stems from the observation of selective destruction of VENs in early stages of fronto-temporal dementia consistent with evidence from functional imaging⁸³.

The involvement of the ACC as a mechanism of alexithymia in later life was suggested by Paradiso et al.⁸⁹ who reported a significant inverse correlation between the grey matter volume of the right ACC and alexithymia, a phenomenon that appeared to be a function of older age. Consistent with this idea, a structural MR imaging study showed that the right pregenual ACC grey matter volume in 7 healthy subjects was negatively correlated with alexithymia⁹⁰, but the small sample size is a limitation. Additional regions showing negative correlations with alexithymia were the right middle and superior temporal gyrus, the right postcentral gyrus, the right precuneus and

the right inferior parietal lobe. Conversely, homologous regions on the left hemisphere were not correlated with alexithymia⁹⁰. Circuitry connecting temporal, parietal and frontal areas has been posited to support processing of information relating to the self and to emotional events⁹¹. Consistent with in vivo structural neuroanatomy studies, functional neuroimaging studies have shown reduced activity of ACC in individuals with high alexithymia scores⁹² and a reduction of grey matter in anterior cingulate cortex and in middle temporal gyrus in healthy women⁷⁸. Paradiso et al.⁹³ recently reported that patients with right middle cerebral artery (MCA) stroke show high levels of alexithymia, perhaps as a consequence of distant functional effects of the stroke damage reducing the functioning of the ACC. The ACC is vascularized by MCA and some parts are highly sensible to hypoxia⁹⁴.

Functional networks involving the frontal lobe

Over the last decade, the concept of the default mode network (DMN) has gained broad acceptance as a set of functionally interconnected brain regions including the dorsal medial and ventral medial prefrontal cortex, the medial and lateral parietal cortex and the temporal cortex and the posterior cingulate cortex⁹⁵⁻⁹⁷. The DMN is active during introspection, while this network is deactivated during “nonself-referential goal-directed tasks in keeping with the folk-psychological notion of losing one’s self in one’s work” (Sheline et al.⁹⁵, p. 1). A recent study showed diminished connectivity within the DMN among participants with alexithymia⁹⁸. The authors studied 20 alexithymic subjects and 18 healthy participants using functional magnetic resonance imaging (fMRI) and observed that brain areas of DMN (medial, frontal, and temporal regions) showed weaker connections in alexithymic subjects than healthy individuals, while connectivity between DMN and sensory-motor areas was higher⁹⁸.

This body of work suggests that alexithymia may be related to malfunctioning of brain structures including the cingulate and prefrontal cortex regulating and subserving emotional awareness⁷⁷ and self-oriented planning, a mental state often associated with emotion. Hence, it is plausible that when mental functions supported by regions in the DMN fail to work properly, the orderly linkage from body stimuli to emotions and to symbolic language is disrupted⁹⁹, leading to poor symbolization of bodily stimuli and alexithymia.

Basal ganglia

Striato-thalamo-cortical circuits support emotional processing¹⁰⁰. Different basal ganglia structures (globus pallidus, ventral striatum, caudate, subthalamic nucleus) have been associated with primary alexithymia⁸⁸ and

with organic alexithymia⁶⁶. Lee et al.¹⁰¹ observed that in 38 healthy subjects with higher levels of alexithymia showed lower activation in right caudate nucleus in response to angry facial stimuli. Six subjects with focal lesions of the left basal ganglia investigated by MRI and PET neuroimaging techniques showed blunted emotions and elevated scores on the assessment of alexithymia¹⁰². High frequency electrical stimulation of the subthalamic nucleus altered emotional perception of (positive or negative) emotional visual stimuli in patients with Parkinson’s disease¹⁰³.

Corpus callosum

In homo sapiens, abstract reasoning, symbolization, language, and introspection are highly developed cognitive functions. Emotional processing requires the capacity to translate body language into symbolic (emotional) language. Some authors observed that patients with agenesis of the corpus callosum showed an impairment in paralinguistic information and prosody and concluded that individuals with, “agenesis of [the] corpus callosum appear to lack interhemispheric integration of critical aspects of language” (Paul et al.¹⁰⁴, p. 1). In an observational study performed on 28 healthy women, language and visuo-spatial abilities were found to be linked to dimensions of the corpus callosum measured by inversion recovery magnetic resonance images. Verbal fluency was found directly correlated with splenium of corpus callosum, whereas an inverse correlation was observed between splenium dimension and language lateralization¹⁰⁵. The importance of the corpus callosum in the pathogenesis of alexithymia has been posited for some time based on the observed dysfunctional transfer of information between right and left hemisphere¹⁰⁶⁻¹⁰⁸.

Frontal lobe

Frontotemporal structures in particular in the right hemisphere play a fundamental role in processing emotions¹⁰⁹ and are involved in frontostriatal circuitry which plays a role in emotional awareness¹¹⁰. The orbitofrontal cortex is widely connected with limbic structures and beyond (i.e., parietal, temporal, occipital)⁶⁹. Moreover, the orbitofrontal cortex is crucial in recognizing emotional vocal and facial expressions^{111 112}.

In a study on 314 alcohol dependent subjects, regression analysis provided evidence that frontal lobe dysfunction, as assessed using the Frontal Systems Behavior Scale, or FrSBe, “mediated the relationship between alexithymia (TAS-20 total score) and risky alcohol use”¹¹³.

The role of the right hemisphere in alexithymia has emerged from studies of patients with right hemisphere stroke^{65 114}. Paradiso et al.¹¹⁵ observed that depressed pa-

tients with right hemisphere stroke often presented with an apathetic and nondysphoric depression believed to be the result of reduced emotional processing abilities. In addition, frontal lobe dysfunction may mediate the relationship between alexithymia and risk for drug addiction¹¹³. The review by Tabibnia and Zaidel¹¹⁶ suggests that the extant data supports the role of deficit in interhemispheric information transmission and poorer right hemisphere functioning in alexithymia. This view was confirmed in a recent (albeit small) study showing that the grey matter of the right hemisphere (not the left) negatively was associated with alexithymia⁹⁰.

Amygdala and insula of Weil

The amygdala is thought to be critical for emotional processing¹¹⁷. It is made up of several nuclei, including the lateral nucleus that connects with sensory cortical areas, the central nucleus that is the output and connects with brain stem areas subserving the neurovegetative manifestations of emotions, and the basal nucleus that is the relay between lateral and central nucleus^{118 119}. The amygdala is a node between the sensory input (to be) labelled with an emotion and the emotional manifestations of the body¹²⁰. Reduced activity in the amygdala can be responsible for an alteration in emotional processing. Because of the key role of amygdala in emotional processing, it is reasonable to speculate that the amygdala can have pathogenetic role in alexithymic subjects.

Difficulty identifying feelings from the faces of others was observed in 21 healthy subjects with a reduced activation of amygdala by fMRI¹²¹. Difficulty identifying emotions was significantly and negatively correlated with the neural response of the amygdala to sad faces¹²¹. Alexithymic patients with anorexia nervosa showed poor amygdala activation (as well as cingulate cortex)¹²².

Higher levels of activation were observed in right insula and in inferior frontal lobe of alexithymic patients¹²³. The insula is connected with the amygdala and ACC and plays an important role as a "prelimbic area" during emotional processing¹²⁴. To the elevated levels of insula activity can be attributed the somatic tendency of individuals with alexithymia.

In summary, several brain regions are purportedly associated with alexithymia. At first glance, alexithymia may appear to be the result of several disparate mechanisms. While this may remain a possibility, a more parsimonious view is that focal dysfunction in differing nodes of emotional processing networks may disrupt the functioning of the entire network. Thus, a pattern of mental and behavioural dispositions consistent with alexithymia may occur in association with dysfunction in differing nodes of the brain supporting emotion processing.

Conclusions

Emotional processing encompasses cortical and subcortical brain mechanisms. Subcortical structures (limbic regions, basal ganglia) may allow for emotional processing to occur at a level below conscious awareness, while the cortex plays a role in (self)-awareness of emotional processing. The interaction between sub-cortical and cortical structures is fundamental for adaptive emotion processing. In the philosophic meditations of ancient Greeks and Romans, emotions were understood as unnecessary passions and were often viewed as *diseases of the mind*. Contemporary views purport that poor recognition and identification of emotions, including personal emotions, may be maladaptive. In this review, we discussed primary alexithymia, understood as a condition arising from various sources including genetic and experiential and distinguished between primary and secondary alexithymia. In addition, we further differentiated secondary alexithymia as deriving either from a psychological stressor or due to an event that indirectly or directly impacts the brain. Thus, we conclude that:

1. recognition and treatment of primary alexithymia is warranted because of its potential as a risk factor for psychiatric disorders⁶⁰. While evidence for a definitive etiological role of alexithymia in physical illness is sparse, alexithymia may exert influence on illness behaviour based on, "physical symptoms, disability, and excessive health care use"¹²⁵;
2. secondary alexithymia may be a response to the psychological distress of an organic disease or psychological trauma;
3. organic alexithymia may be considered to be a specific subtype of secondary alexithymia that is a consequence of brain damage (e.g. traumatic or vascular) to specific regions including the anterior cingulate, basal ganglia, amygdala, insula, right hemisphere, and corpus callosum. In vulnerable individuals with brain injury, alexithymia may increase the risk of suicide⁴².

In future research, it may be important to distinguish between treatment responses in primary, secondary and organic alexithymia³³.

References

- 1 Sifneos PE. *Clinical observations on some patients suffering from psychosomatic diseases*. In: Antonelli F, Ancona L, eds. *Acta psychosomatica*. Roma: SIMP 1967.
- 2 Taylor GJ, Bagby RM, Parker JDA. *Disorders of affect regulation: alexithymia in medical and psychiatric illness*. Cambridge: Cambridge University Press 1997.
- 3 Goleman D. *Emotional intelligence*. London: Bloomsbury 1996.
- 4 Taylor G. *Psychosomatic medicine and contemporary psy-*

- choanalysis. Madison, CT: International Universities Press 1987.
- 5 Krystal, H. *Alexithymia and the effectiveness of psychoanalytic treatment*. IJPP 1982;9:353-78.
- 8 Ogrodniczuk JS, Piper WE, Joyce AS. *Effect of alexithymia on the process and outcome of psychotherapy: a programmatic review*. Psychiatry Res 2011;190:43-8.
- 7 Ogrodniczuk JS, Piper WE, Joyce AS. *The negative effect of alexithymia on the outcome of group therapy for complicated grief: what role might the therapist play?* Compr Psychiatry 2005;46:206-13.
- 8 Nemiah JC, Freyberger H, Sifneos PE. *Alexithymia: a view of the psychosomatic process*. In: Hill O, ed. *Modern trends in psychosomatic medicine*. London-Boston: Butterworth 1976, pp. 430-9.
- 9 Cicero MT. *Tusculan disputations*. New York: Haper & Brothers 1894.
- 10 Damasio AR. *Descartes' error: emotion, reason, and the human brain*. New York: Putnam 1994.
- 11 Goleman D. *Working with emotional intelligence*. London: Bantam Dell Pub Group 2000.
- 12 Salovey P, Mayer JD. *Emotional intelligence*. Imagin Cogn Pers 1990;9:185-211.
- 13 Spencer H. *Principles of psychology*. London: Longman, Brown, Green and Longmans 1855.
- 14 Darwin C. *The expression of emotions in man and animals*. London: John Murray 1872.
- 15 James W. *What is emotion?* Mind 1884;9:188-205.
- 16 Lange CG, James W. *The emotions*. Baltimore: Williams & Wilkins 1922.
- 17 Cannon WB. *Bodily changes in pain, hunger, fear, and rage*. New York: Appleton 1929.
- 18 Arnold MB. *Emotion and personality*. Vol. I. *Psychological aspects*. New York: Columbia University Press 1960.
- 19 Eibl-Eibesfeldt I. *Love and hate: a natural history of behavior patterns (foundations of human behavior)*. London: Methuen 1971.
- 20 Beadle JN, Brown V, Keady B, et al. *Trait empathy as a predictor of individual differences in perceived loneliness*. Psychol Rep 2012;110:3-15.
- 21 Kohler CG, Turner TH, Gur RE, et al. *Recognition of facial emotions in neuropsychiatric disorders*. CNS Spectr 2004;9:267-74.
- 22 Nicolò G, Semerari A, Lysaker PH, et al. *Alexithymia in personality disorders: correlations with symptoms and interpersonal functioning*. Psychiatry Res 2011;190:37-42.
- 23 Messina A, Fogliani AM, Paradiso S. *Association between alexithymia, neuroticism, and social desirability scores among Italian graduate students*. Psychol Rep 2010;107:185-92.
- 24 Lamberty GJ, Holt CS. *Evidence for a verbal deficit in alexithymia*. J Neuropsychiatry Clin Neurosci 1995;7:320-4.
- 25 Henry JD, Phillips LH, Crawford JR, et al. *Theory of mind following traumatic brain injury: the role of emotion recognition and executive dysfunction*. Neuropsychologia 2006;44:1623-8.
- 26 Onor M, Trevisiol M, Spano M, et al. *Alexithymia and aging: a neuropsychological perspective*. J Nerv Ment Dis 2010;198:891-5.
- 27 Paradiso S, Vaidya JG, McCormick LM, et al. *Aging and alexithymia: association with reduced right rostral cingulate volume*. Am J Geriatr Psychiatry 2008;16:760-9.
- 28 Galderisi S, Mancuso F, Mucci A, et al. *Alexithymia and cognitive dysfunctions in patients with panic disorder*. Psychother Psychosom 2008;77:182-8.
- 29 Lane RD, Sechrest L, Riedel R. *Sociodemographic correlates of alexithymia*. Compr Psychiatry 1998;39:377-85.
- 30 Sifneos PE. *Alexithymia, clinical issues, politics and crime*. Psychother Psychosom 2000;69:113-6.
- 31 Mattila AK, Kronholm E, Jula A, et al. *Alexithymia and somatization in general population*. Psychosom Med 2008;70:716-22.
- 32 Lumley MA, Neely LC, Burger AJ. *The assessment of alexithymia in medical settings: implications for understanding and treating health problems*. J Pers Assess 2007;89:230-46.
- 33 Spalletta G, Serra L, Fadda L, et al. *Unawareness of motor impairment and emotions in right hemispheric stroke: a preliminary investigation*. Int J Geriatr Psychiatry 2007;22:1241-6.
- 34 Bermond, B. *Brain and alexithymia*. In: Vingerhoets A, Van Bussel F, Boelhouwer J, eds. *The (non)expression of emotions in health and disease*. Tilburg: Tilburg University Press 1997, pp. 115-29.
- 35 Bailey PE, Henry JD. *Alexithymia, somatization and negative affect in a community sample*. Psychiatry Res 2007;150:13-20.
- 36 Kano M, Hamaguchi T, Itoh M, et al. *Correlation between alexithymia and hypersensitivity to visceral stimulation in human*. Pain 2007;132:252-63.
- 37 Barsky AJ. *Amplification, somatization, and the somatoform disorders*. Psychosomatics 1992;33:28-34.
- 38 Nakao M, Barsky AJ, Kumano H, et al. *Relationship between somatosensory amplification and alexithymia in a Japanese psychosomatic clinic*. Psychosomatics 2002;43:55-60.
- 39 De Gucht V, Heiser W. *Alexithymia and somatisation: quantitative review of the literature*. J Psychosom Res 2003;54:425-34.
- 40 Rasmussen NH, Agerter DC, Colligan RC, et al. *Somatization and alexithymia in patients with high use of medical care and medically unexplained symptoms*. Ment Health Fam Med 2008;5:139-48.
- 41 Jellesma FC, Rieffe C, Terwogt MM, et al. *Do I feel sadness, fear or both? Comparing self-reported alexithymia and emotional task-performance in children with many or few somatic complaints*. Psychol Health 2009;24:881-93.
- 42 Wood RL, Williams C, Lewis R. *Role of alexithymia in suicide ideation after traumatic brain injury*. Psychiatry Res 2011;190:43-8.
- 43 Honkalampi K, Koivumaa-Honkanen H, Tanskanen A, et al. *Why do alexithymic features appear to be stable? A*

- 12-month follow-up study of a general population. *Psychother Psychosom* 2001;70:247-53.
- 44 Taylor GJ, Bagby RM, Parker JD. *Alexithymia. State and trait.* *Psychother Psychosom* 1993;60:211-4.
 - 45 Bach M, de Zwaan M, Ackard D, et al. *Alexithymia: relationship to personality disorders.* *Compr Psychiatry* 1994;35:239-43.
 - 46 McCormick LM, Brumm MC, Beadle JN, et al. *Mirror neuron function, psychosis, and empathy in schizophrenia.* *Psychiatry Res* 2012;201:233-9.
 - 47 Beadle JN, Paradiso S, Salerno A, et al. *Alexithymia, emotional empathy, and self-regulation in anorexia nervosa.* *Ann Clin Psychiatry* 2013;25:107-20.
 - 48 Newton-Howes G, Tyrer P, Johnson T. *Personality disorder and the outcome of depression: meta-analysis of published studies.* *Br J Psychiatry* 2006;188:13-20.
 - 49 Viinamäki H, Hintikka J, Tanskanen A, et al. *Partial remission in major depression: a two-phase, 12-month prospective study.* *Nord J Psychiatry* 2002;56:33-7.
 - 50 Vanheule S, Desmet M, Verhaeghe P, et al. *Alexithymic depression: evidence for a depression subtype?* *Psychother Psychosom* 2007;76:135-6.
 - 51 Gallo JJ, Rabins PV. *Depression without sadness: alternative presentations of depression in late life.* *Am Fam Physician* 1999;60:820-6.
 - 52 Paradiso S, Vaidya J, Tranel D, et al. *Nondysphoric depression following stroke.* *J Neuropsychiatry Clin Neurosci* 2008;20:52-61.
 - 53 Paradiso S, Caspers K, Tranel D, et al. *Cognition and nondysphoric depression among adoptees at high risk for psychopathology.* *Compr Psychiatry* 2011;52:498-506.
 - 54 Freyberger H. *Supportive psychotherapy techniques in primary and secondary alexithymia.* *Psychother Psychosom* 1977;28:337-42.
 - 55 Lesser IM. *A review of the alexithymia concept.* *Psychosom Med* 1981;43:531-43.
 - 56 Krystal H. *Alexithymia and psychotherapy.* *Am J Psychother* 1979;33:17-31.
 - 57 Wearden A, Cook L, Vaughan-Jones J. *Adult attachment, alexithymia, symptom reporting, and health-related coping.* *J Psychosom Res* 2003;55:341-7.
 - 58 Kano M, Mizuno T, Kawano Y, et al. *Serotonin transporter gene promoter polymorphism and alexithymia.* *Neuropsychobiology* 2012;65:76-82.
 - 59 Wise TN, Mann LS, Mitchell JD, et al. *Secondary alexithymia: an empirical validation.* *Compr Psychiatry* 1990;31:284-8.
 - 60 de Vente W, Kamphuis JH, Emmelkamp PM. *Alexithymia, risk factor or consequence of work-related stress?* *Psychother Psychosom* 2006;75:304-11.
 - 61 Yehuda R, Steiner A, Kahana B, et al. *Alexithymia in Holocaust survivors with and without PTSD.* *J Trauma Stress* 1997;10:93-100.
 - 62 Zeitlin SB, McNally RJ, Cassidy KL. *Alexithymia in victims of sexual assault: an effect of repeated traumatization?* *Am J Psychiatry* 1993;150:661-3.
 - 63 Becerra R, Amos A, Jongenelis S. *Organic alexithymia: a study of acquired emotional blindness.* *Brain Inj* 2002;16:633-45.
 - 64 De Panfilis C, Salvatore P, Marchesi C, et al. *Parental bonding and personality disorder: the mediating role of alexithymia.* *J Pers Disord* 2008;22:496-508.
 - 65 Spalletta G, Pasini A, Costa A, et al. *Alexithymic features in stroke: effects of laterality and gender.* *Psychosom Med* 2001;63:944-50.
 - 66 Huang MF, Yeh YC, Tsang HY, et al. *Alexithymia associated with bilateral globus pallidus lesions after carbon monoxide poisoning.* *Kaohsiung J Med Sci* 2010;26:333-6.
 - 67 Ross ED, Homan RW, Buck R. *Differential hemispheric lateralization of primary and social emotions.* *Neuropsychiatry Neuropsychol BehavNeurol* 1994;7:1-19.
 - 68 Adolphs R, Damasio H, Tranel D, et al. *Cortical systems for the recognition of emotion in facial expressions.* *J Neurosci* 1996;16:7678-87.
 - 69 Barbas H. *Connections underlying the synthesis of cognition, memory, and emotion in primate prefrontal cortices. Proceedings of the human cerebral cortex: from gene to structure and function.* *Brain Res Bull* 2000;52:319-30.
 - 70 Koponen S, Taiminen T, Honkalampi K, et al. *Alexithymia after traumatic brain injury: its relation to magnetic resonance imaging findings and psychiatric disorders.* *Psychosom Med* 2005;67:807-12.
 - 71 Vijayaraghavan L, Vaidya JG, Humphreys CT, et al. *Emotional and motivational changes after bilateral lesions of the globus pallidus.* *Neuropsychology* 2008;22:412-8.
 - 72 Messina A, Fogliani AM, Paradiso S. *Alexithymia in oncologic disease: association with cancer invasion and hemoglobin levels.* *Ann Clin Psychiatry* 2011;23:125-30.
 - 73 Ogrodniczuk JS, Sochting I, Piper WE, et al. *A naturalistic study of alexithymia among psychiatric outpatients treated in an integrated group therapy program.* *Hum Psychopharmacol* 2009;24:331-6.
 - 74 Cravello L, Caltagirone C, Spalletta G. *The SNRI venlafaxine improves emotional unawareness in patients with post-stroke depression.* *Brain Lang* 2003;85:313-24.
 - 75 Smith M, Daurat A, Pariente P, et al. *French translation of Schalling-Sifneos Personality Scale Revised and Beth Israel Questionnaire, 2 evaluation tools of alexithymia.* *Encephale* 1992;18:171-4.
 - 76 Bush G, Luu P, Posner MI. *Cognitive and emotional influences in anterior cingulate cortex.* *Trends Cogn Sci* 2000;4:215-22.
 - 77 Gündel H, López-Sala A, Ceballos-Baumann AO, et al. *Alexithymia correlates with the size of the right anterior cingulate.* *Psychosom Med* 2004;66:132-40.
 - 78 Borsci G, Boccardi M, Rossi R, et al. *Alexithymia in healthy women: a brain morphology study.* *J Affect Disord* 2009;114:208-15.
 - 79 Paus T. *Primate anterior cingulate cortex: where motor control, drive and cognition interface.* *Nat Rev Neurosci* 2001;2:417-24.

- 80 Bermond B, Vorst HC, Moormann PP. *Cognitive neuropsychology of alexithymia: implications for personality typology*. Cogn Neuropsychiatry 2006;11:332-60.
- 81 Phillips ML, Drevets WC, Rauch SL, et al. *Neurobiology of emotion perception I: the neural basis of normal emotion perception*. Biol Psychiatry 2003;54:504-14.
- 82 Mohanty A, Engels AS, Herrington JD, et al. *Differential engagement of anterior cingulate cortex subdivisions for cognitive and emotional function*. Psychophysiology 2007;44:343-51.
- 83 Allman JM, Tetreault NA, Hakeem AY, et al. *The von Economo neurons in the fronto-insular and anterior cingulate cortex*. Ann NY Acad Sci 2011;1225:59-71.
- 84 Kaufman JA, Paul LK, Manaye KF, et al. *Selective reduction of Von Economo neuron number in agenesis of the corpus callosum*. Acta Neuropathol 2008;116:479-89.
- 85 Hakeem A, Sherwood C, Bonar C, et al. *Von Economo neurons in the elephant brain*. Anat Rec 2009;292:242-8.
- 86 Stanyon R, Consigliere S, Morescalchi MA. *Cranial capacity in hominid evolution*. Human Evolution 1993;8:205-16.
- 87 Butti C, Sherwood C, Hakeem A, et al. *Total number and volume of Von Economo neurons in the cerebral cortex of cetaceans*. J Comp Neurol 2009;515:243-59.
- 88 Evrard H, Forro T, Logothetis N. *Von Economo neurons in the anterior insula of the macaque monkey*. Neuron 2012;74:482-9.
- 89 Paradiso S, Vaidya JG, McCormick LM, et al. *Aging and alexithymia: association with reduced right rostral cingulate volume*. Am J Geriatr Psychiatry 2008;16:760-9.
- 90 Sturm VE, Levenson RW. *Alexithymia in neurodegenerative disease*. Neurocase 2011;17:242-50.
- 91 Buckner RL, Andrews-Hanna JR, Schacter DL. *The brain's default network: anatomy, function, and relevance to disease*. Ann NY Acad Sci 2008;1124:1-38.
- 92 Berthoz S, Artiges E, Van De Moortele PF, et al. *Effect of impaired recognition and expression of emotions on fronto-cingulate cortices: an fMRI study of men with alexithymia*. Am J Psychiatry 2002;159:961-67.
- 93 Paradiso S, Anderson BM, Boles Ponto LL, et al. *Altered neural activity and emotions following right middle cerebral artery stroke*. J Stroke CerebrovascDis 2011;20:94-104.
- 94 Vaidya JG, Paradiso S, Boles Ponto LL, et al. *Aging, gray matter, and blood flow in the anterior cingulate cortex*. Neuroimage 2007;37:1346-53.
- 95 Sheline YI, Barch DM, Price JL, et al. *The default mode network and self-referential processes in depression*. PNAS 2009;69:113-6.
- 96 Bressler SL, Menon V. *Large-scale brain networks in cognition: emerging methods and principles*. Trends Cogn Sci 2010;14:277-90.
- 97 Zhang L, Zhu C, Ye R, et al. *Impairment of conflict processing in alexithymic individuals*. Neurosci Lett 2011;504:261-4.
- 98 Liemburg EJ, Swart M, Bruggeman R, et al. *Altered resting state connectivity of the default mode network in alexithymia*. Soc Cogn Affect Neurosci 2012;7:660-6.
- 99 Tekin S, Cummings JL. *Frontal-subcortical neuronal circuits and clinical neuropsychiatry: an update*. J Psychosom Res 2002;53:647-54.
- 100 Dondaine T, Péron J. *Emotion and basal ganglia (I): what can we learn from Parkinson's disease?* Rev Neurol 2012;168:634-41.
- 101 Lee BT, Lee HY, Park SA, et al. *Neural substrates of affective face recognition in alexithymia: a functional magnetic resonance imaging study*. Neuropsychobiology 2011;63:119-24.
- 102 Vijayaraghavan L, Adolphs R, Kennedy DP, et al. *A selective role for right insula. Basal ganglia circuits in appetitive stimulus processing*. Soc Cogn Affect Neurosci 2012, in press.
- 103 Brücke C, Kupsch A, Schneider GH, et al. *The subthalamic region is activated during valence-related emotional processing in patients with Parkinson's disease*. Eur J Neurosci 2007;26:767-74.
- 104 Paul LK, Van Lancker-Sidtis D, Schieffer B, et al. *Communicative deficits in agenesis of the corpus callosum: nonliteral language and affective prosody*. Brain Lang 2003;85:313-24.
- 105 Hines M, Chiu L, McAdams LA, et al. *Cognition and the corpus callosum: verbal fluency, visuospatial ability, and language lateralization related to midsagittal surface areas of callosal subregions*. Behav Neurosci 1992;106:3-14.
- 106 Buchanan DC, Waterhouse GJ, West SC. *A proposed neurophysiological basis of alexithymia*. Psychother Psychosom 1980;34:248-55.
- 107 Larsen JK, Brand N, Bermond B, et al. *Cognitive and emotional characteristics of alexithymia: a review of neurobiological studies*. J Psychosom Res 2003;54:533-41.
- 108 Romei V, De Gennaro L, Fratello F, et al. *Interhemispheric transfer deficit in alexithymia: a transcranial magnetic stimulation study*. Psychother Psychosom 2008;77:175-81.
- 109 Phan KL, Wager T, Taylor SF, et al. *Functional neuroanatomy of emotion: a meta-analysis of emotion activation studies in PET and fMRI*. Neuroimage 2002;16:331-48.
- 110 Northoff G, Heinzel A, de Greck M, et al. *Self-referential processing in our brain. A meta-analysis of imaging studies on the self*. Neuroimage 2006;31:440-57.
- 111 Ross ED, Mesulam MM. *Dominant language functions of the right hemisphere: prosody and emotional gesturing*. Arch Neurol 1979;36:144-8.
- 112 Hornak J, Rolls ET, Wade D. *Face and voice expression identification in patients with emotional and behavioural changes following ventral frontal lobe damage*. Neuropsychologia 1996;34:247-61.
- 113 Lyvers M, Onuoha R, Thorberg FA, et al. *Alexithymia in relation to parental alcoholism, everyday frontal lobe functioning and alcohol consumption in a non-clinical sample*. Addict Behav 2012;37:205-10.
- 114 Lumley MA, Sielky K. *Alexithymia, gender, and hemispheric functioning*. Compr Psychiatry 2000;41:352-59.
- 115 Paradiso S, Vaidya J, Tranel D, et al. *Nondysphoric depression*.

- sion following stroke. *J Neuropsychiatry Clin Neurosci* 2008;20:52-61.
- ¹¹⁶ Tabibnia G, Zaidel E. *Alexithymia, interhemispheric transfer, and right hemispheric specialization: a critical review.* *Psychother Psychosom* 2005;74:81-92.
 - ¹¹⁷ Zald DH. *The human amygdala and the emotional evaluation of sensory stimuli.* *Brain Res Rev* 2003;41:88-123.
 - ¹¹⁸ Amunts K, Kedo O, Kindler M, et al. *Cytoarchitectonic mapping of the human amygdala, hippocampal region and entorhinal cortex: intersubject variability and probability maps.* *Anat Embryol* 2005;210:343-52.
 - ¹¹⁹ Kapp BS, Whalen PJ, Supple WF, et al. *Amygdala contributions to conditioned arousal and sensory information processing.* In: Aggleton JP, ed. *The amygdala: neurobiological aspects of emotion, memory, and mental dysfunction.* New York: Wiley-Liss 1992, pp. 229-54.
 - ¹²⁰ LangevinJP. *The amygdala as a target for behavior surgery.* *Surg Neurol Int* 2012;3:S40-6.
 - ¹²¹ Kugel H, Eichmann M, Dannlowski U, et al. *Alexithymic features and automatic amygdala reactivity to facial emotion.* *Neurosci Lett* 2008;435:40-4.
 - ¹²² Miyake Y, Okamoto Y, Onoda K, et al. *Brain activation during the perception of stressful word stimuli concerning interpersonal relationships in anorexia nervosa patients with high degrees of alexithymia in an fMRI paradigm.* *Psychiatry Res* 2012;201:113-9.
 - ¹²³ Moriguchi Y, Decety J, Ohnishi T, et al. *Empathy and judging other's pain: an fMRI study of alexithymia.* *Cereb Cortex* 2007;17:2223-34.
 - ¹²⁴ Mayberg HS. *Limbic-cortical dysregulation: a proposed model of depression.* *J Neuropsychiatry Clin Neurosci* 1997;9:471-81.
 - ¹²⁵ Lumley MA, Stettner L, Wehmer F. *How are alexithymia and physical illness linked? A review and critique of pathways.* *J Psychosom Res* 1996;41:505-18.
 - ¹²⁶ TenHouten WD, Hoppe KD, Bogen JE, et al. *Alexithymia: an experimental study of cerebral commissurotomy patients and normal control subjects.* *Am J Psychiatry* 1986;143:312-6.
 - ¹²⁷ Heinzl A, Minnerop M, Schäfer R, et al. *Alexithymia in healthy young men: a voxel-based morphometric study.* *J Affect Disord* 2012;136:1252-6.
 - ¹²⁸ Parker JD, Keightley ML, Smith CT, et al. *Interhemispheric transfer deficit in alexithymia: an experimental study.* *Psychosom Med* 1999;61:464-8.