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Which patient will feel down, which will be happy? The need to study the genetic disposition of emotional states

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Abstract

Purpose In quality-of-life (QL) research, the genetic susceptibility of negative and positive emotions is frequently ignored, taken for granted, or treated as noise. The objectives are to describe: (1) the major findings of studies addressing the heritable and environmental causes of variation in negative and positive emotional states and (2) the major biological pathways of and genetic variants involved in these emotional states.

Methods Literature overview.

Results The heritability estimates for anxiety and depression are 30–40%. Related traits as neuroticism and loneliness are also highly heritable. The hypothalamo-pituitary-adrenal axis is the ‘final common pathway’ for most depressive symptoms. The many findings of investigated genes are promising but not definitive. Heritability estimates of positive emotional states range between 40 and 50%. Life satisfaction and mental health share common genetic factors with optimism and self-esteem. The prefrontal cortex is a candidate brain area for positive emotional states. Biological and genetic research into positive emotional states is scarce.

Conclusion Genetically informative studies may provide insights into a wide variety of complex questions that traditional QL studies cannot deliver. This insight in turn will help us to design more effective supportive programs that could moderate the outcomes of genetically based predispositions.

Keywords Review · Positive emotional states · Negative emotional states · Biological pathways · Genes
Introduction

Psychological or mental functioning is one of the key components of quality of life (QL) and encompasses the entire spectrum of psychological well-being: from anxiety and depression on the one hand to happiness and life satisfaction on the other. Since being confronted with a disease is one of the major stressors in life, it is understandable that patient-based QL research has focused primarily on the negative end of this range. Frequently used QL questionnaires, such as the European Organisation for Research and Treatment of Cancer QL questionnaire (EORTC QLQ-C30), exclusively inquire about negative emotions (e.g., being tense, worried, irritable, depressed) [1]. In the area of health psychology, the stress process has also been described predominantly in terms of negative emotions. This focus may have stemmed from the acknowledgment that negative emotions have adaptive value as they mobilize the fight/flight response and focus attention on the problem at hand [2].

However, there is mounting evidence that positive emotions co-occur with negative emotions during intensely stressful periods of life [2, 3]. Positive emotions have also been found to have adaptive significance as they may broaden the individual’s attentional focus and behavioral repertoire, thereby increasing the person’s intellectual, social, and physical resources [4]. There is a growing empirical body of research showing that positive emotions have a salutary impact on health, particularly regarding the immune system response and pain tolerance [5, 6]. In other words, happy people are in general healthier. The World Health Organization has started to emphasize happiness as a component of health [7], and the British Medical Journal has recently published an article on the dynamic spread of happiness along connections in a large social network and the implications for healthcare [8].

The balance between negative and positive emotions may differ within persons over time, e.g. immediately after the diagnosis of a disease, negative emotions will likely prevail. The balance between negative and positive emotions also differs across individuals. We all know persons whose view on life is gloomy and somber irrespective of happy and fortunate circumstances. Conversely, we can all name people who persist in a sunny and cheerful view on life despite setbacks and mayhem. In patient-based QL research, this inborn nature is frequently ignored, taken for granted, or treated as noise. The field of QL research would benefit from taking this genetic component into account. Insight into the genetic and environmental components of patient-reported emotional states will ultimately allow us to explore new pathways for improving patient care. If we can identify patients who have a high risk of experiencing poor psychological functioning, we will be able to better target psychological support and/or pharmacological treatment. In other words, we will be able to direct our limited resources in a timely fashion to those who are most likely to need them.

Research on twins, multi-generational families, and population-based cohorts has provided ample empirical evidence of a genetic predisposition for negative emotional states, such as depression, anxiety, and loneliness. Additionally, an increasing number of studies showed substantial heritability of positive emotional states, such as subjective well-being, happiness, and life satisfaction. The objectives of this paper are first to describe the major findings of the studies addressing the heritable and environmental causes of variation in individual differences in negative and positive emotional states. Second, we will describe the major biological pathways of and genetic variants involved in these emotional states. We will thus describe those biological pathways and genetic variants that affect QL, which is consistent with the adapted theoretical model of Wilson and Cleary [9] as described by Sprangers et al. [10]. Since more biological and genetic research has been conducted in negative than in positive emotional states, the paper will reflect this imbalance.

Negative and positive emotional states are commonly reported in separate papers. We purposefully bring these findings together as helping patients to achieve a high level of well-being requires targeting both negative and positive affect.

Negative emotional states

Causes of individual differences

Genetic as well as environmental factors play a role in the etiology of negative emotional states, such as anxiety and depression. The heritability estimates for both anxiety and depression disorders range between 30 and 40%. These percentages most likely represent the lower bound. Actual heritability may be higher when measurement error could be ruled out and depression could be perfectly reliably assessed [11, 12]. The remaining variance can be attributed to individual-specific environmental influences, as
Loneliness over ages 7, 10, and 12 [20]. The longitudinal findings in adults [18]. This finding of a heritability of around 50% is replicated in children, using an average score of older adults [18]. The heritability estimates of neuroticism have also been found to range around 30–40% [17]. Findings in adolescents generally support findings in adults and young children that neuroticism is influenced principally by genetic and unique, non-shared environmental factors [16].

In this section and the next section on genetic variants, we will focus primarily on anxiety and depression, as these negative emotional states elicited a great deal of biological and genetic research. The hypothalamo-pituitary-adrenal (HPA) axis is considered to be the “final common pathway” for most depressive symptoms [13] and thus may be important for patient-reported distress. Indeed, a large part of the environmental and genetic risk factors for depression appear to correlate with increased HPA-axis activity in adults. Figure 1 illustrates the pathogenesis of depression as formulated by Bao et al. [13]. In depressed patients, stress acting on the HPA axis results in a disproportionately high activity of this system due to a deficient cortisol feedback effect. Such an impaired negative feedback mechanism and overreaction of the HPA-axis to stress may be based on genetic factors and environmental factors, including placental dysfunction, smoking of the pregnant mother, or child abuse. High corticotropin-releasing hormone (CRH) and cortisol levels contribute to symptoms of depression by their central effects. The set point of the HPA-axis is at least partly genetically determined. For example, the heritability estimate of basal cortisol levels was found to be 62%, as indicated by a simultaneous analysis of five comparable twin studies [21].

Many patients may also experience the pain of loneliness that a chronic or disabling disease may induce. Loneliness can be described as feelings of social isolation and dissatisfaction with one’s social relationships. There is robust evidence that loneliness plays a pivotal role in emotional states, such as mood, anxiety, anger, pessimism, and dysphoria [18]. Loneliness also has a significant genetic component. In a cross-sectional study of adult twins, the heritability estimate for variation in loneliness scores averaged over items and measurement occasions was found to be around 50% [19]. In a longitudinal analysis of two individual items (“I feel lonely” and “Nobody loves me”), the heritability in young adults (below 35 years) was found to be 77 and 70% respectively and 41 and 54% in older adults [18]. This finding of a heritability of around 50% is replicated in children, using an average score of loneliness over ages 7, 10, and 12 [20]. The longitudinal analyses in this study show, however, that heritability is 58% at age 7, 56% at age 10 but drops to 26% at age 12. A parallel increase in influences of shared family environment is observed, explaining 6% of the variance at age 7, 8% at age 10, and 35% at age 12. The remaining variance is explained by relatively stable influences of non-shared environmental factors. Stability in loneliness is high with correlations over time ranging between 0.51 and 0.69, indicating that individuals who score high on loneliness at a young age have a substantial chance to stay lonely throughout childhood, whereas gregarious youngsters have a small chance of suddenly becoming lonely.

**Biological pathways**

In this section and the next section on genetic variants, we will focus primarily on anxiety and depression, as these negative emotional states elicited a great deal of biological and genetic research. The hypothalamo-pituitary-adrenal (HPA) axis is considered to be the “final common pathway” for most depressive symptoms [13] and thus may be important for patient-reported distress. Indeed, a large part of the environmental and genetic risk factors for depression appear to correlate with increased HPA-axis activity in adults. Figure 1 illustrates the pathogenesis of depression as formulated by Bao et al. [13]. In depressed patients, stress acting on the HPA axis results in a disproportionately high activity of this system due to a deficient cortisol feedback effect. Such an impaired negative feedback mechanism and overreaction of the HPA-axis to stress may be based on genetic factors and environmental factors, including placental dysfunction, smoking of the pregnant mother, or child abuse. High corticotropin-releasing hormone (CRH) and cortisol levels contribute to symptoms of depression by their central effects. The set point of the HPA-axis is at least partly genetically determined. For example, the heritability estimate of basal cortisol levels was found to be 62%, as indicated by a simultaneous analysis of five comparable twin studies [21].
High cortisol levels in depression may result in an impaired dopamine system [22], resulting in anhedonia (insensitiveness to pleasure or incapacity for experiencing happiness). Additionally, decreased levels of serotonin in the brain are thought to be of importance in anxiety disorders [23], panic disorders [24], and mood disorders, although this latter relation is most probably reflecting a vulnerability to suffer from depressive disorders [25].

Furthermore, sex hormone levels and particularly fluctuations in sex hormone levels may play an important role in the vulnerability to mood disorders [13]. Finally, the suprachiasmatic nucleus, i.e. the biological clock, which regulates circadian and circannual variations in neuronal, hormonal, and behavioral activity, is also involved. It is supposed to be related to circadian and circannual fluctuations in mood and to sleeping disturbances in depression and to hyperactivity of the HPA-axis [13].

Genetic variants

The genetic liability of a common familial disorder like major depression involves multiple genes. Potentially important genes have emerged that are related to the HPA axis, e.g., those that affect AVP, CRH, or cortisol synthesis as well as the production of their respective receptors. Another frequently studied gene is catechol-o-methyltransferase (COMT) that is related to the monoamine catabolism. Many other candidate genes have been proposed and investigated.

To date, the genetic underpinning of depression has been basically studied in three ways. First, linkage studies of informative families have been conducted to identify chromosomal regions (loci) likely to contain genes that contribute to susceptibility of depression. For example, several genome-wide linkage studies have identified regions of chromosomes 15q [26–28], 17p, and 8p [27] to be related to depression. Second, candidate gene association studies have focused on functional polymorphisms (DNA sequence variations that alter the expression and/or functioning of the gene product) in previously identified and new loci encoding for potentially relevant genes, as exemplified above. Many findings of this candidate gene approach can be considered promising but not definitive. The multiple genes involved in depression each exerts a small effect, making genetic linkage and association studies challenging [29]. Some consistent patterns have only recently emerged. For example, the following five genes were significantly associated with major depressive disorder in meta-analyses of polymorphisms that had been investigated in at least three studies [30]: apolipoprotein E (APOE), guanine nucleotide-binding protein (GNB3), methylenetetrahydrofolate reductase (MTHFR), dopamine transporter (SLC6A3), and serotonin transporter (SLC6A4).

Third, genome-wide association studies examine many polymorphisms simultaneously in large samples of unrelated, population-based cases (those with depression) and controls (those without depression). The first genome-wide association study of depression examined more than 430,000 single nucleotide polymorphisms (SNPs) in 1,738 cases of major depression and 1,802 controls and suggested preliminary evidence for the involvement of the presynaptic protein piccolo (PCLO) on chromosome 7 [31]. The results from multiple replication cohorts (6,079 independent cases with major depressive disorder and 5,893 controls) remained inconclusive. However, reanalysis of the PCLO replication study indicated that there was convincing evidence for the potentially causal association of major depressive disorder with one particular SNP, rs2522833, in PCLO [32]. Interestingly, the second genome-wide association study based on two large, independent data sets and a further combined analysis using a meta-analytical approach [33] failed to identify any SNP that achieved significance. The authors concluded that SNPs with substantial odds ratio are unlikely to exist for major depression disorder, at least among the studied SNPs to date.

A consistent, negative finding is worth mentioning. According to the monoamine-deficiency hypothesis, deficits in serotonin were thought to play a predominant role in the pathophysiology of depression [34]. However, antidepressant drugs made to replenish the lack of these neurotransmitters, like serotonin reuptake inhibitors (SSRIs), were shown not to be effective in large groups of patients suffering from depression [22]. In addition, there is no simple relationship between serotonin levels in the brain and mood [25]. Moreover, the fact that SSRIs take weeks to become effective makes also clear that the serotoninergic system cannot play a primary role in mood disorders. Finally, a recent meta-analysis was conducted of the interaction between the serotonin transporter gene (5-HTTLPR) and stressful life events on depression using a total sample of 14,250 participants divided into cases and controls. This study did not yield evidence that this serotonin transporter gene alone or in interaction with stressful life events is associated with an elevated risk of depression [35].

Positive emotional states

Causes of individual differences

An increasing number of twin studies showed substantial heritability of positive emotional states, such as subjective well-being and life satisfaction. Heritability estimates ranged between 40 and 50%, whereas the remaining
Biological pathways and genetic variants

Several studies suggest that the prefrontal cortex is a candidate brain area for happiness and positive emotional states that may be related to taste [53], smell [54] or other inputs via the somatosensory system [55]. Some electroencephalographic (EEG) studies suggest that positive emotional states are associated with increased left cortical power in the alpha frequency compared to the right hemisphere [56, 57]. There is also evidence that dopamine modulates positive emotional states [58], indicating a role for the ventral tegmental area. At the subcortical level, a number of neuropeptide systems have been implicated in positive emotional states, e.g., neotensin and cocaine- and amphetamine-regulated transcript (CART) (both closely associated with dopamine), neuropeptide Y, and oxytocin [58]. Finally, reduced activity of the neuroendocrine [59–61] and cardiovascular systems [58], as well as increased activity of the immune system [58], may all be involved in positive emotional states. However, genes or genomic regions of interest for positive emotional states have not been published at the time of paper writing.

Discussion

We have described the major results of the studies addressing the heritability of and biological pathways and genetic variants involved in negative and positive emotional states, respectively. These affect states are to a substantial degree heritable, with positive states being slightly more heritable (40–50%) than negative states (30–40%). The remaining variances can be attributed to environmental influences unique to the individual, in which the intrauterine period may play an important role. Perhaps contrary to some people’s expectation, emotional states are not affected by environmental influences shared by family members. Another finding is that these mood states share common genetic factors with related personality traits, such as neuroticism for negative emotional states and optimism and self-esteem for positive emotional states. An abundance of studies have focused on the delineation of the biological pathways of negative affect, with the HPA axis considered as the ‘final common pathway’ of depressive symptoms. By contrast, biological and genetic research into positive emotional states is scarce. Multiple genes are involved in emotional states, each exerting a small effect. While the rate of progress is dazzling, particularly for negative affect, the biological complexities do not allow definitive answers yet.

A note of caution is warranted. The studies reviewed employed very different samples, study designs, measures, and analyses thereby potentially limiting their comparability.
Particularly, the highly divergent self-report measures across studies merit attention as reviews and meta-analyses generally ignore these different sources. The question arises whether such collapsing of data is warranted. The cited study by Bar-tels and Boomsma [45] examining the etiology of four different operationalizations of positive affect is enlightening. They found that the four measures were explained by one underlying genetic factor and genetic factors specific to the four measures were negligible. The implication of this finding is that distinct measures of subjective well-being are not distinct at the genetic level and represent biological overlapping constructs. Thus, different studies that collect distinct measures of positive emotional states can be compared. Moreover, pooling data across measures is acceptable, at least for these four measures of positive emotional state. Whereas this finding is heartening, further research is needed to examine the extent to which different measures assessing similar emotional states, or QL domains, also share a similar genetic factor.

An intriguing and pressing question is the extent to which negative and positive emotional states share the same biological and genetic substrate. There is evidence of some common biological mechanism. For example, the dopamine system is involved in negative as well as positive affect. However, the relatively recent finding that high levels of distress can co-occur with high levels of well-being plead for, at least partly, independent biological mechanisms. Clearly, further studies are needed to disentangle the biological and genetic substrates of negative and positive affect, using data sets that include information on both phenotypes.

To avoid a possible misunderstanding about the implications of heritability studies, we would like to call attention to the fact that a high level of heritability does not mean that environmental influences are unimportant. Genes only influence phenotypes within an environment [46]. For example, experimental manipulations of loneliness were found to have powerful effects on mood, shyness, anxiety, and self-esteem [in 18]. Moreover, interventions based on simple and popular concepts such as committing acts of kindness, expressing gratitude or forgiveness, and thoughtful self-reflection had the power to induce a sustainable increase in levels of happiness [62].

At this point, QL researchers may wonder why we need to know the genes if we can examine the presence and severity of depression, extent, and level of happiness, as well as the related personality characteristics by self-report? Molecular genetic studies may provide insights into a wide variety of complex questions that traditional QL studies cannot deliver. For example, how can we identify patients who are vulnerable to long-lasting distress following a diagnosis? How can we predict which patients will suffer from mood disturbances when taking a specific chemotherapeutic regimen? Why do some pharmacotherapeutic treatments not work in all patients with the ‘same’ level of distress? Which patients would benefit from simple interventions to sustainably increase happiness? Why are some psychotherapies not effective in comparable patients with the same ‘problem’? Why do supportive interventions, which one would expect to benefit all patients, often help only a few? Delineation of biological pathways through which various genetic predispositions propel people toward negative or away from positive emotions is needed. Moreover, insight into which patients will respond to which interventions can only be provided by studies assessing the patients’ relative risk for negative and likelihood of positive emotions using family and molecular genetic approaches in combination with assessments of the risk and protective environmental factors. This insight in turn will help us to design more effective supportive programs that could moderate the outcomes of genetically based predispositions.

For example, if we can identify which patients are genetically predisposed to experience intense negative or extreme positive affect, preventive measures such as self-help programs or cognitive behavioral therapy can be implemented. Further, if the patient is entering into a phase of life where unavoidable stressors are present, such as a diagnosis of a chronic disease, supportive care interventions targeted at negative and positive affect or pharmacotherapy can be used concurrently with other treatments that can ultimately lead to better treatment outcomes, including survival. The individual, genetic profile would thus help indicating which treatment or support would be most helpful for a particular patient.

In sum, the combination of the prognostic value of genetic, environmental risk and protective factors ultimately enables appropriate and effective support [63]. Clearly, molecular genetic testing is a long way from being implemented in clinical settings, and to some, the previously sketched vision might seem science fiction. However, to realize this vision, we believe that QL research should embrace the study of the genetic and environmental influences of both negative and positive affects as the resulting insight will ultimately help our patients to become not only happier but also healthier.

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Appendix: glossary [cited from 64]

Chromosome: Structure that is composed mainly of chromatin, which contains DNA and resides in the nucleus of cells.

DNA (deoxyribonucleic acid): The-stranded molecule that encodes genetic information.

Family study: Assessing the resemblance between genetically related parents and offspring and between siblings living together. Resemblance can be due to heredity or to shared family environment.

Gene: The basic unit of inheritance. A sequence of DNA bases that codes for a particular product.

Genome: All the DNA sequences of an organism.

Genotype: The genetic constitution of an individual.

Heritability: The proportion of the phenotypic differences that can be attributed to genetic differences in a particular population.

Linkage: Close proximity of loci on a chromosome.

Linkage analysis: A technique that detects linkage between DNA markers and traits used to map genes to chromosomes.

Locus (plural, loci): The site of a specific gene on a chromosome.

Mapping: Linkage of DNA markers to a chromosome and to specific regions of chromosomes.

Nucleus: The part of the cell that contains chromosomes.

Phenotype: An observed characteristic of an individual that results from the combined effects of genotype and environment.

Polymorphism: A locus with two or more alleles (alternative forms of a gene at a locus). (Functional polymorphism: DNA sequence variations that alter the expression and/or functioning of the gene product)

Single nucleotide polymorphism (SNP): Sequences in the genome that differ by a single nucleotide between one portion of the population and another.

Twin study: Comparing the resemblance of identical and fraternal twins to estimate genetic and environmental components of variance.

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