Multi-scale characterisation of pasta

Effects of raw materials on water absorption, water distribution, and microstructure

THOMAS STEGLICH

Department of Chemistry and Chemical Engineering
CHALMERS UNIVERSITY OF TECHNOLOGY
Gothenburg, Sweden 2015
Multi-scale characterisation of pasta
Effects of raw materials on water absorption, water distribution, and microstructure
THOMAS STEGLICH
ISBN 978-91-7597-152-0

©THOMAS STEGLICH. 2015

Doktorsavhandlingar vid Chalmers tekniska högskola. Ny serie
Nr 3833
ISSN 0346-718X

Department of Chemistry and Chemical Engineering
Chalmers University of Technology
SE-412 96 Gothenburg
Sweden
Telephone + 46 (0)31-772 1000

Cover:
Cross section of cooked spaghetti visualised by Magnetic Resonance Imaging and Light Microscopy. The figure illustrates radial changes in water-macromolecule interactions as well as in microstructure.

Dixa AB
Gothenburg, Sweden 2015
Multi-scale characterisation of pasta
Effects of raw materials on water absorption, water distribution, and microstructure
THOMAS STEGLICH
Department of Chemistry and Chemical Engineering
Chalmers University of Technology

ABSTRACT
Pasta is a product with a long history, but is also still being developed today. Producers want to use new raw materials to make pasta more nutritious, less allergenic, and less dependent on durum wheat. All have in common that new raw materials shall not compromise the desired texture properties of cooked pasta such as the “al dente” feeling.

To facilitate the development of new pasta products, understanding the microstructure of pasta can be a tool. Water transforms and interacts with the microstructure during cooking and the outcome determines the texture. The main objective of this work was to analyse the interplay of microstructure and water, and how this is affected by the choice of raw materials.

We combined light microscopy and Magnetic resonance imaging (MRI) to study the microstructure and water distribution of pasta. We improved the resolution of MRI to yield data in 3D and were able to link MRI data to microstructure components such as fibre particles and the extent of starch gelatinisation.

Monitoring microstructure transformations during cooking and warm-holding of pasta revealed that some transformations are not dependent on the raw materials used. Water ingress towards the core is regulated by starch gelatinisation, which holds true both during cooking and warm-holding. The extent of the continuous starch and protein transformation from core to surface in cooked pasta is mainly governed by the product geometry. Also, texture changes during warm-holding depended mainly on the amount of available water within and around the pasta after cooking.

Nevertheless, raw materials are of importance: A higher protein content limited the degree of starch swelling within the gelatinised region in cooked pasta. Fibre particles can hinder water migration locally due to their perpendicular alignment against the direction of water ingress. Bran particles in particular do not absorb water during cooking, but redistribute it around the particles and create a strong variation in the degree of starch swelling. The severity of this effect correlated with bran particle size.

This thesis provides a comprehensive overview over how local microstructure and raw material choice affect water distribution at different scales in the cooked product.

Keywords: pasta, light microscopy, magnetic resonance imaging, microstructure, water distribution, texture properties, starch, gluten, bran, dietary fibre
List of Publications

This thesis is based on the work contained in the following papers, referred to by Roman numerals in the text:

I  Microstructure and water distribution of commercial pasta studied by microscopy and 3D magnetic resonance imaging  

II Multi-scale characterization of pasta during cooking using microscopy and real-time magnetic resonance imaging  
   D. Bernin1, T. Steglich1, M. Röding, A. Moldin, D. Topgaard, M. Langton  
   1Authors contributed equally  

III Bran particle size influence on pasta microstructure, water distribution, and sensory properties  
   T. Steglich, D. Bernin, A. Moldin, D. Topgaard, M. Langton  
   Submitted to Cereal Chemistry

IV Texture of pasta during warm-holding  
   T. Steglich, A. Moldin, M. Langton  
   Manuscript, intended for publication in Journal of Texture Studies

Contribution report*

I Planned, performed and evaluated the experiments (except for MRI parameter estimation), primarily responsible for writing and revising the manuscript

II Planned, performed and evaluated the experiments, primarily responsible for writing and revising the manuscript

III Planned, performed and evaluated the experiments, and wrote major parts of the manuscript

IV Planned, performed and evaluated the experiments, and wrote the manuscript

* MRI experiments in paper I-III were planned, developed, performed and evaluated in close collaboration with Diana Bernin. All other types of experiments were performed by the author
**List of abbreviations**

BFLM  Bright-field light microscopy
MRI   Magnetic Resonance Imaging
OCT   Optimal cooking time
PLM   Polarized light microscopy

**List of sample abbreviations**

*Industrial-scale spaghetti*

DS     Durum semolina
DS+FB  Durum semolina and wheat fibre
DS+WG  Durum semolina and durum whole-wheat flour
DS+SW  Durum semolina and soft wheat flour

*Laboratory-scale spaghetti*

S40D60 40% starch powder, 60% durum wheat flour
D100   100% durum wheat flour
G20D80 20% gluten powder, 80% durum wheat flour
G40D60 40% gluten powder, 60% durum wheat flour
WW0    0% durum whole-wheat flour, 100% durum wheat flour
WW50   50% durum whole-wheat flour, 50% durum wheat flour
WW100  100% durum whole-wheat flour, 0% durum wheat flour
S440   10% bran fraction (median particle size 440 µm), 90% durum wheat flour
S370   10% bran fraction (median particle size 370 µm), 90% durum wheat flour
S160   10% bran fraction (median particle size 160 µm), 90% durum wheat flour
S90    10% bran fraction (median particle size 90 µm), 90% durum wheat flour

D100 and WW0 were produced with different batches of the same flour type.
# Table of Contents

Introduction ........................................................................................................................................ 1
Objectives ......................................................................................................................................... 3
Background ....................................................................................................................................... 4
   Durum wheat components .............................................................................................................. 4
      Starch ........................................................................................................................................ 4
      Protein/Gluten .......................................................................................................................... 6
      Bran/non-starch polysaccharides .............................................................................................. 7
Pasta processing ................................................................................................................................. 7
   Durum wheat pre-processing ......................................................................................................... 7
   Mixing .......................................................................................................................................... 7
   Extrusion ...................................................................................................................................... 8
   Drying .......................................................................................................................................... 8
Transformations during cooking ........................................................................................................ 9
Water transport models .................................................................................................................... 10
Influence of starch and gluten properties on pasta quality ............................................................ 11
Influence of bran on pasta quality .................................................................................................. 12
Transformations after cooking: Storing pasta ................................................................................ 12
Materials and Methods .................................................................................................................. 13
   Spaghetti preparation .................................................................................................................. 13
   Cooking and Warm-holding ........................................................................................................ 14
   Water absorption ....................................................................................................................... 14
   Light microscopy ....................................................................................................................... 14
   Magnetic resonance imaging .................................................................................................... 15
   Cooking quality ......................................................................................................................... 15
   Statistical analysis .................................................................................................................... 15
Magnetic resonance imaging of pasta: Challenges and method development ......................... 16
   Basic principles of MRI .............................................................................................................. 16
   Challenges in MRI for pasta research ...................................................................................... 17
   MRI in this work ......................................................................................................................... 17
      MRI of cooked pasta .............................................................................................................. 17
      Real-time MRI during cooking of pasta .................................................................................. 18
      Combining MRI and light microscopy ................................................................................... 19
Results and Discussions ................................................................................................................ 21
   General structural transformations ............................................................................................. 21
      Cooking: Water absorption and structure transformations ................................................... 21
      Microstructure of cooked spaghetti ....................................................................................... 23
      Post-cooking: Warm-holding .................................................................................................... 24
      Post-cooking: Storing at ambient temperatures ..................................................................... 26
      Effects of raw materials on texture, microstructure, and water distribution in pasta .......... 27
         Starch and protein content .................................................................................................. 27
         Bran addition and bran particle size ................................................................................... 32
Conclusions ....................................................................................................................................... 39
Future research ................................................................................................................................. 40
Acknowledgements ......................................................................................................................... 41
References ....................................................................................................................................... 42
Introduction

Pasta is a universal food mainly made from wheat, but also from rice and other cereals. It has a century-old history with roots in China and Italy. However, the industrial production did not start before the 1950s (De Vita, 2009). Even in Italy the consumption of pasta rose first after this time, from being limited to feast days to everyday use. Today, Italians consume about 25 kg per capita and year. For comparison: Swedes consume 9 kg per capita and year (International Pasta Organisation, 2012).

Pasta and noodles are offered in manifold forms and ways - as fresh pasta, instant pasta and, most importantly, dried pasta. However, it is still not completely understood how the textural properties of cooked pasta are formed and influenced by every step of the production chain - starting from the choice of raw materials over production itself to end with how to keep the product warm after cooking.

A better understanding of these processes can aid to develop and improve pasta products. The challenges are plenty: nowadays, cooked pasta is often stored before consumption; e.g. by holding it warm in a canteen kitchen or keeping it cold in ready-to-eat meals and salads. Other pasta products are produced with high amounts of dietary fibre to improve their nutritional profile, but this affects texture properties.

Studying pasta microstructure and its interactions with water can be one tool to improve the understanding of the material. Already more than 30 years ago, Resmini and Pagani (1983) noted:

“Pasta proves to be an interesting limited water-starch-protein system where starch/protein competition for water, conformational changes and mutual interactions take place during processing and cooking. The understanding of these phenomena, which may parallel that of other cereal products, can be enhanced by the study of pasta fine structure”.

The focus of the current research was to combine microstructure analysis with a spatially resolved analysis of water distribution in pasta and relate the findings to texture properties.
Objectives

The goal of this work was to better understand how raw materials affect in cooked pasta texture properties such as firmness and stickiness. The main objective was therefore to characterise the interplay of water and microstructure during and after cooking of pasta.

A schematic overview shows the focus of the individual papers (Figure 1).

The specific aims were:

- Improve the spatial resolution of MRI to determine water distribution in pasta after cooking in detail (I, III)
- Improve the temporal resolution of MRI to monitor water migration in pasta during cooking (II)
- Determine the interplay of water and microstructure of pasta during cooking (II)
- After cooking (I, III)
- After warm-holding (IV)
- Elucidate the effects of raw materials (determined by their variation in protein, starch and fibre/bran content) on water absorption, water distribution, and microstructure as well as texture properties (I-IV)
  - Determine in detail the effects of bran particle size (III)
- Define the factors causing texture changes in warm-held pasta (IV)

Figure 1 Overview of main analytical focus of each respective study
Background

The term pasta describes generally sheeted or extruded wheat dough products. They are often categorised into (Asian) noodles and pasta (Marchylo et al., 2004). While noodles are based on common wheat flour and are sheeted, pasta is mainly based on durum wheat flour and is extruded.

The ultimate goal for every pasta manufacturer is to produce pasta with the best texture properties possible. What defines the best cooking quality might be subjective and consumer preference can vary from country to country, but cooking quality is often linked to being high firmness, low stickiness as well as overcooking tolerance (Marti et al., 2014).

Recently, much research has been directed to changing the raw materials used while maintaining the cooking properties by adapting the production process. The research efforts can be grouped into three trends (references refer to reviews):

- Substitute durum wheat with cheaper and local crops (Fuad and Prabhasankar, 2010),
- Increase nutritional value such as content of dietary fibre (Rawat and Indrani, 2014; Sissons and Fellows, 2014), and
- Decrease allergenicity (in particular replace gluten) (Hager et al., 2012; Marti et al., 2014; Petitot et al., 2009).

Durum wheat semolina is seen as the most suitable raw material for pasta production. As a starting point for any raw material modulation, it is useful to understand how the major durum wheat components starch and protein form the structure for desired texture properties during cooking. Texture properties are related to microstructural changes during cooking, which in turn are affected by water and temperature.

Durum wheat components

For pasta production, generally only parts of the wheat grain are used, while whole-wheat pasta requires using the whole grain (Manthey and Schorno, 2002). The main parts of a grain are the endosperm, bran and germ (Figure 2). The endosperm consists primarily of starch granules and protein bodies, which are grouped in a cellular structure surrounded by thin cell walls (Kill and Turnbull, 2001). The bran comprises several protective cell layers while germ is the embryo of the grain and thus is rich in vitamins, minerals, antioxidants and dietary fibre (Manthey and Schorno, 2002).

To achieve homogeneous flours, only the endosperm is used and thus bran and germ are often removed during milling. The remaining endosperm is milled to flour and for common and durum wheat these flours are called farina and semolina, respectively. The texture affecting wheat components starch, protein and bran will be described in the following section.

Starch

Starch is composed of the two polysaccharides amylose and amylopectin, which both are based on glucose residues. Amylose consists of a linear chain with $\alpha$-1,4 linkages, while amylopectin is highly branched through additional $\alpha$-1,6 linkages. The ratio between amylose and amylopectin can vary, but is generally one to three in wheat (Delcour et al., 2010). Amylose and amylopectin are synthesized in granular form with alternating amorphous and semi-crystalline growth rings. The semi-crystallinity makes starch granules birefringent and visible in polarised light (Delcour et al., 2010).
Starch granules are formed during biological synthesis together with non-starch molecules such as phosphates and lipids (Conde-Petit, 2003; Tang et al., 2006). In wheat they show a bimodal distribution, with larger oval type-A-granules and smaller round type-B-granules (Soh et al., 2006).

In the research of this work, also soft wheat flour has been used. Therefore, durum and soft wheat starch are briefly compared. Durum wheat starch has a higher water binding capacity, slightly higher amylose content and lower gelatinisation temperature than common wheat starch (Delcour et al., 2010; Vansteelandt and Delcour, 1999). Zweifel (2001) concluded that this suggests a less compact starch granule structure in durum wheat. However, durum wheat lacks the protein puroindoline which induces a stronger interaction of proteins with the starch granule surface and makes the kernels very hard (Delcour et al., 2010). This may be the reason that during milling starch damage is higher in durum than in common wheat. Compared to common wheat, especially the type-A-granules of durum wheat are smaller (13-16 µm compared to 22-36 µm) and the volume share of them is higher (Pérez and Bertoft, 2010; Soh et al., 2006; Wilson et al., 2006). Finally, according to Vansteelandt and Delcour (1999), durum wheat starch and common wheat starch show about the same protein content, but differ in lipid content.

When sufficient moisture is available, native starch gelatinises when heated above the gelatinisation temperature, losing its crystallinity and structural organization (Copeland et al., 2009). At the same time there is a significant increase in molecular water mobility (Cuq et al., 2003). Water is absorbed first to the amorphous zones and, if there are channels, water first
reaches the inside and then diffuses outward (Copeland et al., 2009; Fannon et al., 2004; Langton and Hermansson, 1989). This hydration process can be quite fast, with a half time of 7s (Lemke et al., 2004). The loosely bound amylose molecules start to leach out of the granule. Starch molecules can swell to a certain amount before they are disrupted by the induced stress. Observing the melting process of starch using differential scanning calorimetry (DSC) measurements, an additional peak appears at higher temperatures than the peak corresponding to starch gelatinisation. This is attributed to the melting of amylose-lipid complexes (Conde-Petit, 2003).

**Protein/Gluten**

The heterogeneous wheat protein is classified according to extractability into four fractions: water soluble albumins, globulin (soluble in salt solution), gliadins (soluble in ethanol) and glutenins (soluble in dilute acids, Troccoli et al., 2000). Albumins and globulins represent the minor part (15-20 %) and have emulsifying properties. The major part of wheat proteins consists of gliadin and glutenin which form the composite gluten. Gluten is an important factor in determining pasta quality as it can form a viscoelastic network (Delcour et al., 2012).

Glutenin is further separated into high and low molecular weight subunits. Both subunits are bound together by intermolecular disulphide bonds and form the huge polymer glutenin in a helical structure (Sissons, 2008; Zweifel, 2001).

In the dry grain the gluten is accumulated in protein bodies which are unevenly distributed in the kernel, both quantitatively and qualitatively (Gil-Humanes et al., 2011; Tosi et al., 2011). Protein concentration is higher close to the bran layer and lower in the central starchy endosperm cells. Some protein bodies can even form a continuous matrix enclosing starch granules (Tosi et al., 2011).

Through the addition of water, gluten bodies transform from the crystalline, glassy state to become rubbery, elastic and form a network through inter-molecular bonds (Kill and Turnbull, 2001). Already at a water content of 15 %, the glass transition occurs below room temperature and hence gluten is in the rubber state at the conditions used for pasta dough preparation. At temperatures above 60 °C, again as a function of moisture, hydrated gluten forms three-dimensional aggregates through the establishment of covalent bonds (protein-protein cross links; Sissons, 2008). This thermosetting is irreversible (Cuq et al., 2003). Gluten bond formation both during processing and cooking is described in more detail elsewhere (Bock and Seetharaman, 2012; Wagner et al., 2011).

Kontogiorgos (2011) suggests a new hierarchical model for the structure of the hydrated gluten. At molecular level individual glutenins and gliadins are interacting via various physical and covalent bonds. At the nanoscale a continuous phase is created where gluten polymers form sheets. These resulting sheets can be seen as the building block of the gluten network. The side-by-side arrangement of the sheets gives a nanoporous structure with “confined” water entrapped into the sheets and interacting strongly with the gluten matrix and “bulk” water surrounding it. Between the sheets nanocapillaries are formed. Depending on the packing of the sheets, a three-dimensional network is formed at the microscale. Finally, the gluten network on the macroscale can show various morphologies.

Regarding its rheological properties, glutenin is responsible for elastic properties in a dough, while gliadin acts as a plasticizer, being responsible for viscous properties such as dough extensibility (Sissons et al., 2007). The term gluten strength is a measure for the balance between the two gluten subfractions, i.e. between viscosity and elasticity (Sissons, 2008). Gluten strength can be measured with the gluten index method and a higher gluten index indicates stronger gluten network.
**Bran/non-starch polysaccharides**

Bran is built up of several layers – from the outermost pericarp, to intermediate layers of testa, nucellar and hyaline layer to the protein-rich aleurone layer closest to the endosperm (Figure 2). The structure of each layer has been shown in detail by Surget and Barron (2005). Wheat bran is both rich in dietary fibre as well as protein, as it is composed of about 48% dietary fibre, 16% starch, 18% proteins, 5% fat, 5% sugars and 6% ash (Meuser, 2008).

Other non-starch polysaccharides can be found in the germ and in the cell walls of the endosperm (Sissons, 2008). The cell walls consist of cellulose, lignin, β-glucan and hemicellulose, where arabinoxylans are a major component within hemicellulose (BeMiller, 2010).

Generally, non-starch polysaccharides are often classified as dietary fibres and sorted into soluble and insoluble fibres (BeMiller, 2010). Bran is mostly insoluble while other added fibres such as inulin and guar gum are soluble (Rakhesh et al., 2015).

**Pasta processing**

After the necessary pre-processing of wheat, and milling to semolina or flour, pasta is produced by three main steps: Semolina is polymerised during mixing, the material is compacted and formed by means of sheeting or extruding and this structure is stabilised by drying (Kratzer, 2007) (Figure 3).

**Durum wheat pre-processing**

The aim of the milling process is to separate the endosperm from the germ and the bran of the wheat kernel to get a maximum of semolina at the best quality (Kill and Turnbull, 2001). Milling comprises therefore the following steps: Cleaning of the wheat, tempering (or debranning) to loosen the bran layers as well as milling, and purifying to achieve semolina of the right particle size.

The milling process is a combination of grinding, sifting, and blending to reduce milling passages. To grind pre-processed kernels, they are led over several rolls which are categorised according to their roughness into breakers, reduction rolls, steel screens and purifiers (Cubadda et al., 2009). The breakers open the kernels and separate the carbohydrate-rich endosperm from bran and germ. The reduction rolls are mainly used to adjust the particle size whereas a larger number of purifiers is used to remove remaining bran particles (Delcour et al., 2010).

**Mixing**

Commonly, semolina particles and water are pre-mixed at high-speed to ensure a homogeneous particle wetting (Kill and Turnbull, 2001). Afterwards the mixture is kneaded for some minutes at lower speed to form a dough and especially to form the gluten network. A narrow particle size distribution ensures a homogeneous dough and reduces the risk of the formation of non-wetted particles (white spots) (Kill and Turnbull, 2001). A smaller particle size is preferred to reduce the mixing time. Semolina with a particle size below 250 µm can be mixed in 5 min compared to the coarse semolina which needs 15 min to mix (Rubin, 2007). An alternative, integrated mixing system was introduced some years ago which uses co-rotating twin-screws, leading to intensified mixing and kneading. Thereby, mixing time was reduced to 20 sec (Kill and Turnbull, 2001).

Vacuum in the mixing zone prevents oxidation of semolina natural pigments and the intrusion of air bubbles, which gives a better shine of the pasta product (Pagani et al., 2007). Additionally, temperature should be controlled during dough preparation with an optimum temperature between 35 and 40°C (Kill and Turnbull, 2001).
Extrusion

The formed dough is moved under kneading towards the extrusion zone. The dough is compacted and pressure builds up. The pressure depends on dough moisture and temperature as well as on die resistance which in turn depends on the extrusion area and speed (Kill and Turnbull, 2001).

Extrusion can increase the water solubility of dietary fibre and solubility increases with higher specific mechanical energies (Robin et al., 2012). However, stickiness also increases with higher energy input during extrusion (Kratzer, 2007). Thus, the energy input should be just high enough to form a homogeneous pasta dough (Kratzer, 2007).

Temperature should be kept below 50°C to prevent gluten aggregation already in the extruder as this network would be destroyed by the high shear forces at the die. In general, shear forces (e.g. from worn dies) should be kept at a minimum, as they increase damage of starch granules which can swell excessively during cooking (Petitot et al., 2009).

Commonly two types of dies are available: Bronze and Teflon. While Teflon dies produce a smooth texture and even surface, the surface of pasta extruded through bronze dies is rough. The effect was quantified by Lucisano et al. (2008). The bronze die extruded spaghetti had a higher porosity and breaking strength of the dried spaghetti was reduced by 20-30%. The type of drying cycle and semolina particle size distribution had an influence as well, but to a minor extent.

Drying

Drying reduces the moisture content from roughly 30% to below 12.5% to make pasta a stable product (Pagani et al., 2007). The local water concentration in the pasta differs during drying, which can induce internal stress, potentially leading to fractures and cracks (Migliori et al., 2005). Drying is therefore seen as the critical processing step. Zhang et al. (2013) showed...
that spaghetti shrank faster in radial than in axial direction during drying in an experimental drying chamber. Additionally, the spaghetti shrank torsional, non-central directed in the beginning and linear, central directed later. These findings may explain the arising of internal stress and cracks.

High temperature drying can improve product quality and compensate the use of raw materials of inferior quality to some extent (Petitot et al., 2009; Zweifel et al., 2003). In addition, high temperature drying can reduce overall drying time (Petitot et al., 2009). Zweifel et al. (2003) analysed various drying profiles and concluded that drying at high temperature at a late stage gave the best product quality (compared to low-temperature- and early high-temperature drying). The protein network was preserved through promoting protein denaturation to a dense and continuous network encapsulating starch granules and thus swelling of starch granules was reduced. Compared to early high-temperature drying, late high-temperature drying stabilised the protein network to such an extent that the network was still visible even in the external zone of overcooked pasta.

High temperature drying can make up for inferior protein composition (Del Nobile et al., 2003b). However, if the temperature is too high and the polymerisation reaches a point where the proteins get too rigid to expand, it will result in inferior pasta as well (Delcour et al., 2010). High temperature drying also increases the risk for heat damages of the dried pasta - mainly off-colours and off-flavours and reduced nutritional value of the proteins with the breakdown of lysine due to the formation of Maillard reaction products (de Noni and Pagani, 2010; Peressini, 2011). It may also affect the allergenicity of the proteins (Petitot et al., 2009).

Raw material choice can also affect drying behaviour. Bran inclusion modifies the drying kinetics and the equilibrium moisture is different in bran-rich than in bran-free semolina during drying (Villeneuve and Gelinans, 2007).

**Transformations during cooking**

In their native state and at room temperature, both starch and gluten are in a glassy state and are organised in crystalline structure. They are very limited in their water uptake, with a decreasing solubility for the starch components from amylopectin, to amylose and amylose-lipid complexes (Conde-Petit, 2003). Depending on temperature and water content, starch and gluten can change from a glassy state to a rubbery state reversibly via a so-called glass transition. A second, irreversible transformation occurs at higher temperatures: Starch gelatinisation and protein denaturation.

During pasta cooking, starch granules absorb water and swell. This increases the volume and pressure on the protein network (Delcour et al., 2010). With increasing temperature, two endothermic transitions take place. First, starch gelatinises, and at a higher temperature amyllose-lipid complexes dissociate (Petitot and Micard, 2010). Starch gelatinisation and gluten polymerisation occur at the same time and are competitive in regard to the absorbed water as well as controlled by the water penetration inside the pasta (Petitot et al., 2009).

It is interesting to note that while starch becomes soluble during gelatinisation, gluten becomes insoluble during the network formation (Pagani et al., 2007). Therefore, the kinetics of these processes determines the extent of starch swelling.

Depending on the amount of starch swelling, amylose can leach out of the granules and the starch granules disintegrate. This can induce excessive cooking loss (loss of material into the cooking water) and increased stickiness (Delcour et al., 2010). The final product quality depends on how the protein structure withstands the swelling of the starch (Delcour et al., 2000a, 2000b). When protein polymerisation during cooking does not make up for insufficient polymerisation during drying, excessive cooking losses occur. In contrast, the proteins in pasta
dried at very high temperatures may well have polymerised to such an extent that they are too rigid to expand and retain the gelatinised starch during cooking, resulting in inferior quality pasta (Bruneel et al., 2010; Delcour and Hoseney, 2009). The latter depends on the protein network, with a strong network preventing the leakage and dissolving of starch granules (Zweifel et al., 2003; Bruneel et al., 2010). Honeycomb-like structure at the surface of partly and fully cooked spaghetti could be detected (Heneen and Brismar, 2003; Sung and Stone, 2005). In any case, the cooking generates a concentrical change from the core to the surface in the microstructure (Cunin et al., 1997). This gradient in the change of microstructure and moisture content with a firm core is often referred to as ‘al dente’. Delcour et al. (2010) summarised the cooking process by arguing that the transformation of starch is a hydration-driven gelatinisation process in the outer layer while it is a heat-induced crystallite melting in the centre of the pasta.

**Water transport models**

Water is a major part of most foods and its distribution steers food properties. In the context of pasta products, especially the water absorption and desorption processes are of interest. During the life-time of a pasta product, water migration occurs during mixing of the dough, drying and cooking of the pasta, and finally during storage of the cooked pasta. A lot of research has been carried out to model these sorption processes. The modelling is complicated by the fact that the sorption is happening at the same time as swelling/shrinking and physical modifications of the structural elements, i.e. gelatinisation of starch and denaturation of gluten (Del Nobile et al., 2003). To ease the mathematical description of these phenomena, water migration is most often described in one dimension only. That is also why spaghetti and lasagne plates are used so abundantly, as the radial profile of the spaghetti and the large surface compared to the thickness of lasagne, respectively, leads to a water migration in predominantly just one dimension.

Numerous models have been reported that tried to explain the water absorption process with models based on stationery as well as non-stationary Fickian diffusion and it is further disputed whether dried pasta can be seen as porous medium or not. To mention just one rather new study: Zhu et al. (2011) developed a finite element analysis taking into account the simultaneous effects of water absorption and viscoelastic deformation. They describe pasta as an unsaturated food system from the beginning. The moisture transport during the hydration process is the sum of diffusion and viscoelastic effects and thus, the water transport is dominated by non-Fickian diffusion.

The most complete model may have been proposed by Del Nobile et al. (2003b). According to them, the hydration process occurring during cooking and overcooking depends on four main phenomena: 1. melting kinetics of the crystalline starch domain, 2. water diffusion, 3. relaxation of the macromolecular matrix (swelling), and 4. residual deformation release. Water diffusion is controlling water uptake at the beginning of the rehydration process. The diffusion occurs through two phenomena at the same time: A. Molecular diffusion (related to Brownian movement), driven by the concentration gradient of water (low in the core of the pasta and high at the surface). B. Macromolecular matrix relaxation, driven by the disequilibrium of the local system. The second effect means that when water hits the macromolecular matrix (that is starch and protein), the matrix does not take up all the water needed to reach its equilibrium immediately. The matrix swells over time and the kinetics of the swelling is dependent on the level of disequilibrium. Melting of starch crystals requires a minimum temperature, but is then a fast process compared to the processes mentioned before. The fourth phenomenon (release of residual deformation) takes into account that, during drying, the continuous protein phase is
“frozen” into a state which is not in equilibrium. During hydration and melting of the starch crystals, the reformation can occur which reduces the macromolecular matrix. However the fourth phenomenon is outweighed by the amount of water taken up, which leads to an increase in spaghetti size during the rehydration process.

In a further study by the same authors, a moisture profile over the radius of a cooked spaghetti was modelled which indicates that first during starch melting the matrix takes up significant amount of water (Del Nobile et al., 2003a).

All the models so far concentrated mostly on the geometrical aspect of the hydration process. The direct influence of certain raw materials on the pasta hydration process, however, has not been modelled yet. Del Nobile et al. (2003b) used lab-scale spaghetti made with different wheat varieties, but could not relate any difference to the properties of the raw materials used.

It seems that for so far the influence of raw materials on the pasta hydration process is described only qualitatively and based solely on empirical research. Studies which relate water transport to structural elements (such as starch, gluten, or bran) could not be found.

**Influence of starch and gluten properties on pasta quality**

To understand the influence of the wheat components on pasta quality, several authors carried out reconstitution studies. That means, they fractioned the semolina into gluten (or even several gluten fractions) and starch (variation in starch granule distribution) and combined the fractions in varying compositions together (Delcour et al., 2000; Delcour et al., 2000; Sissons et al., 2002). Sissons et al. (2007) altered the gluten composition and showed that glutenin increased and gliadin decreased dough strength in a dough made of both reconstituted flour and semolina base. HMW-GS increased the dough strength of the base, while LMW-GS decreased it. The dough strength changes did not alter spaghetti texture, however.

An increased share of 32-44 % B-granules resulted in an improved pasta quality, with an increased firmness, reduced stickiness and reduced cooking loss (Soh et al., 2006). The dough strength decreased above a share of 32 % B-granules. The authors speculated, that too many B-granules would need too much water creating imbalanced water distribution throughout the dough.

Increased amylose concentrations led to more extensible dough, lower water uptake of the pasta and increased firmness (Soh et al., 2006). According to the authors, higher amounts of amylose lead to more tightly packed starch granules, which could be more resistant to deformation under swelling. They concluded that the decreased water uptake might change the sensory perception negatively. In line with these observations were findings of inferior cooking properties for reduced amylose content (Gianibelli et al., 2005; Vignaux et al., 2005). An explanation could be that starch granules of low amylose content deteriorate physically more easily and during cooling they form aggregates, but no network. This would result in a soft structure (Tan et al., 2006).

The reconstitution method in itself has some limitations, because fractioning changes material properties. Sissons et al. (2002) showed that the dough strength increased in a reconstituted pasta sample compared with a non-reconstituted sample.

Another approach was used by Wood (2009). Chickpea-fortified spaghetti was produced and compared with standard durum wheat spaghetti to study the mechanisms for pasta quality. Firmness is influenced more by the composition and content of gluten than of protein content in total. Additionally, the protein-polysaccharide matrix rather than the starch composition is important for cooking loss, while cooking loss and stickiness do not necessarily correlate. Finally, increased protein and amylose contents decreased pasta stickiness.
Influence of bran on pasta quality

Due to its nutritional profile, there is a great interest to learn more about how wheat bran and other fibre sources affect pasta properties. Only recently, several groups studied bran and bran fractions being incorporated into pasta/noodles ((Aravind et al., 2012; Chen et al., 2011; Chillo et al., 2008; Kaur et al., 2012; Shiau et al., 2012; West et al., 2013). Bran often induces strong aromas in pasta (due to phenolic acids) and changes texture with increased cooking loss and decreased firmness (West, 2012). What induces these texture changes is not fully understood. Tudorica et al. (2002) argue for fresh pasta that soluble fibres are included into the protein network of the pasta, while insoluble fibres such as bran are disrupting the network. In the case of dried pasta, SEM images showed that the bran particles where not in contact with the protein matrix and thus disrupted the network (Manthey and Schorno, 2002). Others argue that it might depend on the amount and type of fibre (germ particles affected the protein network to a larger extent than bran particles; Aravind et al. 2012) and on the process conditions (Villeneuve and Gelinas, 2007). One recent study rejects the theory of gluten network destruction for the case of bread (Noort et al., 2010). Instead, the authors argue that the fibres interact physically or chemically with the gluten and thus hinder gluten aggregation.

Several approaches have been reported to counteract the deteriorating effect of bran inclusion. Heat treated bran (Sudha et al., 2011), and pasta dried at high temperatures (West et al., 2013) helped to remain the desired texture.

Transformations after cooking: Storing pasta

Storing cooked pasta deteriorates its sensorial properties over time (McCarthy et al., 2002; Olivera and Salvadori, 2011; Wood, 2009). Internal moisture migration has been pointed out as factor for pasta stored at ambient conditions. Moisture redistributes from the outer to the inner zone during storage and reduces both moisture gradient as well as firmness of pasta (Gonzalez et al., 2000). While the moisture content becomes more homogeneous throughout the gelatinised region, the area of the ungelatinised core will remain the same during storage in ambient temperatures (Horigane et al., 2006; Sekiyama et al., 2012).

The deterioration effect can be controlled by the storage process. Irie et al. (2004) treated cooked spaghetti samples differently after production: samples were dried, frozen, or stored at ambient or chilled conditions. Dried and frozen spaghetti showed a clear moisture gradient from surface to core with a low moisture content in the core. Fresh and chilled spaghetti showed a gentle moisture gradient whereas one week stored spaghetti did barely show any gradient. Mechanical properties followed the tendencies in the moisture gradient, with higher forces needed to break dried and frozen spaghetti and lower forces for the chilled spaghetti. Faster product freezing can remain better initial quality during frozen storage (Olivera and Salvadori, 2011).

The choice of raw materials can also influence the properties after storage. Chickpea-fortified spaghetti remained firmer than semolina spaghetti when stored (Wood, 2009). Furthermore, spaghetti maintained firmer after cooking when seasoned with sodium chloride or monosodium glutamate as those salts temporarily absorbed excess water at the surface of the spaghetti (Horigane et al., 2009).

Warm-holding pasta is, compared to cold-storing, even more challenging for foodservices with pasta becoming softer and less chewy (Al-Obaidy et al., 1984). Texture deteriorates faster in warm-held starch-rich foods compared to cold-storing as starch gelatinisation is continuing (Briffaz et al., 2014a). Trends in texture changes are similar for warm-held and overcooked pasta (Willbrandt et al., 1989). Others found also larger differences in cooking quality as cooking time was increased over optimal cooking time and attributed it more to gluten strength than protein content (Dexter et al., 1981; Grzybowski and Donnelly, 1979).
Materials and Methods

The spaghetti form was chosen for its simple dimensions resulting in one-dimensional, radial water ingress. Analysed were both spaghetti produced on industrial-scale (paper I, IV) as well as laboratory-scale (paper II, III). The industrial-scale spaghetti were based mainly on fine durum semolina (*Triticum durum*, DS), but differed in protein content and quality as the flour of various batches and durum wheat varieties was used (paper I and IV). The product DS+SW contained also soft wheat flour (*Triticum aestivum*). Samples in paper I and IV were produced in different years. The spaghetti in paper I differed furthermore in fibre content and type of fibre. Lab-scale spaghetti of greatly varied protein and starch contents were achieved by mixing durum semolina with either starch or gluten powder (paper II). The effect of bran particle size was studied by milling coarse bran to median particle sizes between 90 and 440 µm and mixing it with durum wheat flour. Additionally, durum wheat flour was mixed with durum whole-wheat flour to vary the whole-wheat content in spaghetti from 0 to 100% (paper III).

The samples studied are listed with an approximated protein content of the respective flours used (Table 1). More compositional data is given in the respective papers.

**Table 1** List of spaghetti samples studied. Protein refers to calculated protein content of the flour mixes (14% wet base).

<table>
<thead>
<tr>
<th>Paper I/IV</th>
<th>Industrial-scale</th>
<th>Protein</th>
<th>Lab-scale</th>
<th>Protein</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paper I</td>
<td>DS</td>
<td>Durum wheat flour</td>
<td>13.6</td>
<td>S40D60</td>
</tr>
<tr>
<td></td>
<td>DS+FB</td>
<td>D. + wheat fiber</td>
<td>12.7</td>
<td>D100</td>
</tr>
<tr>
<td></td>
<td>DS+WG</td>
<td>D. + whole-wheat flour</td>
<td>13.3</td>
<td>G20D80</td>
</tr>
<tr>
<td></td>
<td>DS+SW</td>
<td>D. + soft wheat flour</td>
<td>12.0</td>
<td>G40D60</td>
</tr>
<tr>
<td>Paper IV</td>
<td>DS+SW</td>
<td>D. + soft wheat flour</td>
<td>14.2</td>
<td>S440</td>
</tr>
<tr>
<td></td>
<td>DS1</td>
<td>Durum wheat flour</td>
<td>14.2</td>
<td>S370</td>
</tr>
<tr>
<td></td>
<td>DS2</td>
<td>Durum wheat flour</td>
<td>13.7</td>
<td>S160</td>
</tr>
<tr>
<td></td>
<td>DS3</td>
<td>Durum wheat flour</td>
<td>13.6</td>
<td>S90</td>
</tr>
<tr>
<td></td>
<td>DS+FB*</td>
<td>D. + wheat fiber</td>
<td>13.8</td>
<td>WW0</td>
</tr>
<tr>
<td></td>
<td>DS+WG*</td>
<td>D. + whole-wheat flour</td>
<td>15.1</td>
<td>WW50</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>WW100</td>
<td></td>
</tr>
</tbody>
</table>

* Samples were analysed, but not included in paper IV

**Spaghetti preparation**

All industrial-scale spaghetti were manufactured with the same process and supplied by Lantmännens Cerealia (Järna, Sweden). The lab-scale process was supposed to mimic the industrial process. Due to equipment restriction, lab-scale spaghetti production differed from industrial-scale: Mixing and extrusion was carried out in a small pasta machine (Edelweiss TR/75C, Italy) without vacuum. Spaghetti were extruded at a lower pressure compared to an industrial process and rested outside the oven at ambient temperature for a short term. Finally, lab-scale spaghetti was dried in a combi steamer oven (CCM, Rational, Germany) with limited temperature and humidity control. Diameters of the uncooked spaghetti were 1.65-1.70 mm (paper I and IV), 1.50-1.75 mm (paper II) and 1.55 mm (paper III).
Cooking and Warm-holding

In general, when spaghetti should be analysed in its cooked state, spaghetti was cooked in boiling, distilled water and cooled directly afterwards in ice water. For sensory analysis and warm-holding, however, spaghetti was cooked in salted tap water (0.7% w/w NaCl) and spaghetti in paper II was also cooked in salted distilled water (0.7% w/w NaCl) corresponding to a salt level recommended for sensory evaluation (Delcour et al., 2000b).

The following procedure was established for warm-holding (paper IV). 1 kg spaghetti was batch cooked for 7 or 10 min in 10 L water, drained and transferred in a suitable solid steel pan. The pan was covered and placed in a combi-steam oven set at 80°C and 30% relative humidity to ensure that the spaghetti did not dry during warm-holding. Samples were removed from the centre of the pan after 15 min as well as 90 min of warm-holding. Additionally after 90 min, spaghetti strands were taken from the bottom of the pan (referred to as 90 min bottom).

The phrase optimal cooking time (OCT) will be used in the results sections. This refers to a standardised method (AACC, 2000) and defines the cooking time as the time when the non-gelatinised core disappears.

Water absorption

Determining the weight increase over cooking time is an easy measure for the water distribution at the macroscale. Individual spaghetti strands (25 mm) were cooked and removed in intervals from the water. Directly after cooling in ice water, they were blotted and weighed. The weight increase was determined as the mass ratio between the cooked and the dry sample \((W-W_0)/(W_0)\). Whenever possible, the diameter of each dry spaghetti strand was measured before cooking. Thereby the water absorption data could be normalized with the initial surface area which is determined by the dry diameter. The normalisation followed Ogawa et al. (2011).

Light microscopy

Light microscopy is one of the oldest analytical methods as its working principle is rather simple. A light beam is sent through a set of lenses and the sample to receive an enlarged image of the sample structure on a detector. The image is formed by several sources of contrast such as absorption, scattering and reflection of the illumination. Diffraction limits the image resolution theoretically at about the half of the wave length of the incident light beam - using visible light this gives a limit of about 200 nm. This limit is in practice in the magnitude of \(\mu m\) due to sample preparation and properties. Thus, light microscopy enables to analyse food products from the micro- to the macroscale (Autio and Salmenkallio-Marttila, 2001). The main reason to use light microscopy was its ability to map all important components starch, gluten and bran/fibre in one image.

After cooking, pasta samples were prepared for sectioning either by plastic embedding (paper I) or freezing (paper II-IV). The former allows thinner sections of 1 \(\mu m\) (improving the resolution), but involves long embedding procedures which may affect the microstructure. Furthermore, in undercooked pasta the plastic embedding medium did not diffuse into the ungelatinised core which made it impossible to slice whole cross-sections.

Freezing is much faster and easier to produce longitudinal sections with, at the cost of having thicker sections of 5-10 \(\mu m\). Additionally, slices do not attach on the specimen holder as well as plastic sections. This might have introduced artefacts during staining especially in bran-rich pasta.

Sections were analysed both in polarized light (paper I-IV) as well as bright-field light (paper I-III). Only structures that show birefringence are visible in polarised light. This includes ungelatinised starch as well as fibre structures which makes polarised light useful to distinguish the cooking front as well as bran and cell-wall material within the starch-gluten matrix.
In bright-field light microscopy, contrast is enhanced by staining agents. Light Green stains proteins while iodine stains starch. In the resulting images, proteins are stained in green, starch granules in blue/violet, amylose in blue and amylopectin in brown. Fibre structures are generally not stained (but appeared often greenish in micrographs of bran-rich spaghetti which was likely due to remaining staining agent light green). Thus, in the stained light micrographs, all important components within pasta could be visualised in the same image. The structures can be seen in the magnified light micrographs in Figure 5. In the core region, starch appeared as dark blue granules (small and large granules) surrounded by gluten network in green. In the intermediate region, swollen starch granules appeared more violet while amylose in dark blue accumulated in the core and surfaces of starch granules. In the surface region, starch phase separated into amylose enrichments (blue) and random, brownish structures of amylopectin. The image of the intermediate region includes an aleurone cell rich in protein (green), fat (brown) and is surrounded by cell walls (cellulose, unstained).

**Magnetic resonance imaging**

Magnetic resonance imaging can be used to map the water content and water-macromolecule interactions within pasta. Some studies reported using MRI in pasta research, but we still needed to adapt the technique to fulfil our demands.

The basic principles and challenges of MRI are therefore discussed in a separate chapter, together with how we addressed these challenges and how we finally used MRI in this work.

**Cooking quality**

Cooking quality include properties defining pasta in its cooked, eatable state, such as appearance, texture and flavour. In this work some texture properties were determined instrumentally as well as sensorially.

Breaking strength of uncooked and firmness as well as stickiness of cooked and warm-held spaghetti were determined instrumentally (Instron Universal Testing Machine Model No. 5542, 500 N load cell; Instron, High Wycombe, UK) following reported routines (AACC, 2000; Fiorda et al., 2013; Shiau et al., 2012; Sissons et al., 2008). Samples were cooked in excess of water for 10 min (paper I and IV) or 8 min (paper III) and cooled for 1 min in ice water for firmness measurement, while they were only drained for stickiness measurement. Firmness was determined as maximum compression force and stickiness as maximum retraction force; both determined in N.

For sensory analysis (paper III), bran-rich spaghetti was cooked for 10 min instead of 8 min as pre-test showed that 8 min cooked spaghetti were experienced as undercooked. Firmness, stickiness, elasticity, surface roughness and whole-wheat flavour were evaluated by 8 semi-trained panellists in a difference from control test following Bustos et al. (2011a). The panel assessed each sample twice by determining how much more or less intense a certain attribute was compared to the reference WW50. Attributes were assessed on a linear scale (-5 to +5 with the reference placed in the centre at 0). Additionally, the panel was asked to rate each spaghetti for overall liking (0 to 10 without a reference).

**Statistical analysis**

Texture and sensory analysis data were subjected to one-way analysis of variance (ANOVA, General linear model). Grouping information was obtained using the Tukey Method with a 95.0% confidence interval. P-values less than 0.05 were considered significant. Analyses were performed using Minitab 16 Statistical Software (Minitab Inc., USA).
Magnetic resonance imaging of pasta: Challenges and method development

Cooked pasta contains roughly about 60% water and it is thus of interest how water affects and interacts with other components such as starch, gluten and fibre. $^1$H magnetic resonance imaging (MRI) is a non-invasive method to visualise interactions of water with its surroundings and its distribution in pasta (Lai and Hwang, 2004). In this chapter, the technique will be briefly explained as well as how MRI has been applied to pasta research. Furthermore, for pasta, MRI caused some measurement challenges and it will be described how we addressed these challenges. The chapter is concluded by describing how we combined MRI and light microscopy.

Basic principles of MRI

MRI can be used to measure concentration of compounds, structural elements, temperature or diffusivity in various materials (Mariette et al., 2009). MRI uses the magnetic moment of nuclei – in this work the hydrogen proton (which is abundant in water and organic compounds). The protons in a strong magnetic field will align with the field (Schmidt, 2007). A combination of radio frequency (r.f.) pulses, which are part of a pulse sequence, is then used to excite the protons. The signal is acquired before the spins relax back to the thermal equilibrium (alignment with the field). Multidimensional images e.g. 3D can be obtained by using magnetic field gradients which are orthogonal to each other (Cabrer et al., 2005).

The signal intensity in each voxel depends on the physiochemical environment and is a product of the proton density $I_0$ and the corresponding attenuation factors e.g. relaxation times $T_1$ and $T_2$ (Cabrer et al., 2005). The longitudinal relaxation time $T_1$ is driven by thermal relaxation. The influence of $T_1$ on the signal intensity can be minimised by adjusting two MRI parameters: long repetition times (the time lapse between acquiring and re-exciting the signal should be at least five times $T_1$) and small flip angles of the excitation pulse (Cabrer et al., 2005; Schmidt, 2007).

If $T_1$-weighting can be neglected, the equation for signal intensity $S(TE)$ as a function of the echo time ($TE$) can be shortened to

$$S(TE) = I_0 \exp\left(-\frac{TE}{T_2}\right),$$

where $I_0$ represents the initial intensity or proton density and $T_2$ represents the transverse relaxation time. By running a series of echo times, $T_2$ and $I_0$ can be estimated for each voxel based on a regression of the signal intensities on the aforementioned equation.

$I_0$ can be used as a proxy for water content as mainly water protons are measured. Exchangeable protons and mobile protons from the macromolecules are measured as well, but their share is rather low (see detailed discussion in paper I). $T_2$ is the weighted average of all proton pools (water, starch, gluten, fibre). We interpret therefore $T_2$ as describing the water-macromolecule interactions, which is mainly chemical exchange.

Several factors determine the resolution of an image where some factors are defined by the MRI magnet itself while others can be chosen by the operator. Generally, stronger magnetic field gradients and smaller radiofrequency coils (smaller coil diameter and thus smaller field-of-view) will improve the spatial resolution of a Fourier-transformed image. Furthermore, the shorter the echo time can be, the higher the signal intensity will be. Factors such as the number of voxels and field-of-view as well as number of signal accumulations are a tradeoff between spatial and temporal resolution and the signal-to-noise ratio. I.e. a smaller voxel size results in a decreased signal-to-noise ratio and thus a noisier image.
Challenges in MRI for pasta research

In the last fifteen years, only few research groups have used MRI to study the water distribution and interaction in pasta and noodles. Texture properties were linked to the water distribution in cooked lasagna (Gonzalez et al., 2000; McCarthy et al., 2002), while Bonomi et al. (2012) correlated sensory perception and protein network changes in spaghetti. Lai and Hwang, (2004) studied changes in water interaction during storage at ambient temperatures. Most of the research, however, was carried out in Japan studying both noodles and spaghetti (Horigane et al., 2009, 2006; Irie et al., 2004; Kojima et al., 2004, 2001; Maeda et al., 2009; Sekiyama et al., 2012).

In most studies two-dimensional images (slices) were acquired to achieve a high temporal resolution. This approach allows a high in-plane spatial 2D resolution (Bonomi et al. (2012) reported $35 \times 35 \mu m^2$), but averages the signal from a slice at least 1 mm in thickness. Thus the signal cannot be connected to any of the structural elements starch, gluten and fibre individually. Valuable information has been extracted from these measurements, but the level of detail remains limited.

Most of the aforementioned studies report only $T_2$ based data, except for Gonzalez et al. (2000) and McCarthy et al. (2002) who presented signal intensity images. Some studies suggested that the moisture content can be determined by $T_2$ with the help of a calibration curve (Horigane et al., 2006; Irie et al., 2004; Maeda et al., 2009). Pulverised durum semolina was mixed with known amounts of water and cooked until full starch gelatinisation. Then the $T_2$ value of the gelatinised sample was measured and correlated with the known moisture content. This approach neglects, however, that starch in cooked pasta exists in various degrees of gelatinisation depending on the position within the pasta (Heneen and Brismar, 2003), which should influence $T_2$ (Ritota et al., 2008). Additionally, also gluten changes its confirmation during cooking which should influence $T_2$ as well. At least egg protein decreased $T_2$ during its denaturation at constant moisture between 40 and 80°C before increasing during further heating (Mariette, 2009).

Thus, the challenges for this work were to find a suitable setup to acquire MRI signal in 3D at high spatial resolution and to find a more comprehensive interpretation of $I_0$ and $T_2^*$ in the pasta context.

MRI in this work

In paper I-III, we extracted both $I_0$ as well as $T_2$ parameter maps from the measured signal. We studied cooked pasta (paper I and III) as well as pasta during cooking (paper II). The former allowed having a high spatial resolution and a low temporal resolution, while the latter required a high temporal resolution, which reduced the spatial resolution of the MRI measurement.

MRI of cooked pasta

One aim of this work was to improve the spatial resolution of the MRI measurement of cooked spaghetti to be better able to relate the MRI data to structural components within pasta such as fibre particles (paper I and III). The resolution was set to $78 \times 78 \times 78 \mu m^3$ and was measured over the whole spaghetti volume enabling us to visualise the MRI data in several directions (Figure 5). Due to the high resolution and low moisture content, special care was taken to treat the noise (detailed description in paper I).

The chosen resolution resulted in a comparatively long measurement time of 92 min. Such a long time was deemed unsuitable by others as water diffusion and homogenization will take place during the measurement (Maeda et al., 2009). The homogenization effect has been shown by several studies, but despite of changes in absolute values for $I_0$ and $T_2$, the relative differences remained (Horigane et al., 2006; Lai and Hwang, 2004; McCarthy et al., 2002; Sekiyama et al., 2012). The presented $I_0$ and $T_2$ parameter maps may not show the situation directly after cooking, but are still sufficient for comparing various pasta samples being measured with the same protocol.
To minimise the measurement time, we applied a gradient-echo pulse sequence. The gradient-echo uses a 30° excitation pulse, which reduces the repetition time, but also yields the parameter $T_2^*$ (paper I and III). A true $T_2$ can only be determined when a spin echo (90° r.f. pulse and a train of 180° r.f. pulses) is used, such as in the RARE (Rapid Acquisition Relaxation Enhancement) pulse sequence (paper II). $T_2^*$ will not refocus the inhomogeneity of the applied r.f. pulses in contrast to $T_2$.

Measurement time was further reduced from 92 min to 16 min in paper III. We adjusted the field-of-view to acquire the signal only from a representative slab in the centre of the spaghetti instead of the whole volume (Figure 4 A). This could be done because cooked spaghetti shows an isotropic water distribution throughout the cross section (Horigane et al., 2006).

A note concerning the resolution: The ultimate goal would be to achieve a resolution similar to light microscopy. Currently, this would give a very low signal-to-noise ratio or very long measurement times and would require many repetitions. Still, resolution improvements are likely possible. We tested an about ten times higher resolution with a measurement time of 112 min instead of 16 min (Figure 4 B). At this resolution, a quantitative analysis was not possible because the noise became prevalent. However, the exemplary image revealed more details as areas of low or no signal were easier to distinguish from the rest of the pasta matrix. Still, this illustrates potential for higher resolution.

![Figure 4](image-url) $T_2$ parameter maps showing sketches of varied MRI parameters: measured volume (A) and resolution (B). Slices in (B) were obtained from the same sample (S160).

**Real-time MRI during cooking of pasta**

We adapted the MRI method of Mohorič et al. (2004) who studied the cooking process of rice kernels. While rice kernels cook in about 30 min, the cooking process of spaghetti to optimal cooking takes place within 10 min. Hence, real-time MRI during pasta cooking requires the highest temporal resolution possible. A test showed that 3D real-time MRI was possible in principle (resolution 156×156×156 µm$^3$), but resulted in a measurement time of 60s per image. For the data presented in paper II, we chose instead a 2D real-time MRI set-up (156×156×1000 µm$^3$) and one image was recorded in 31s. This greatly reduced moisture migration during the time lapse of an experiment.

Another challenge was to acquire the signal from the spaghetti only and not from the surrounding cooking water. By using a repetition time of 0.9s, the cooking water was heavily $T_1$-weighted and resulted in strongly decreased signal intensity. However, a $T_1$ analysis at the end of the cooking process showed that the signal within the spaghetti was also slightly $T_1$-weighted and thus the uncorrected $I_0$ parameter maps reflected not the true water concentration.
As $T_1$-weighting of the signal varied between the samples, a comparison of $I_0$ between the samples was not possible. Still, the $T_1$-weighted $I_0$ parameter maps showed the temporal changes of the water distribution throughout the cooking series in a qualitative manner.

**Combining MRI and light microscopy**

Acquiring MRI data in 3D allows visualising the water distribution not only in higher resolution, but allows plotting 2D slices in various directions (Figure 5). This is especially helpful when analysing components creating an anisotropic pasta structure such as fibre. Even though MRI data can be plotted in 3D, data will be presented in the results chapter as 2D slices. We experienced that interpreting the data is easier in 2D compared to 3D.

MRI acquires the signal from the whole voxel volume which had a side length of 78 $\mu$m. To illustrate the size of the base area of a voxel, three light micrographs are shown in Figure 5. The images show the voxel volume comprises more starch granules in the core region than in the intermediate or surface region. Bran particles will often be positioned in adjacent voxels as exemplary shown with part of an aleurone cell in the image of the intermediate region.

Spaghetti sections can be cut similarly to the MRI 2D plots in axial, longitudinal and tangential direction and thereby light microscopy analysis can be correlated to MRI data.

The symbols for the three directions shown in Figure 5 will be used throughout this work.

---

**Figure 5** 2D $T_2^*$ parameter maps selected out of a full 3D MRI data set are shown surrounded by a spaghetti surface rendering. 2D plots show three different directions. To illustrate the voxel size of the MRI data, light micrographs are shown in the size of the base area of one voxel. For comparison, thin sections corresponding to the MRI plots are shown in light microscopy (BFLM and PLM). Very dark areas in light micrographs are folded sections. The images show as an example bran-rich spaghetti (DS+WG and S370). Symbols indicate viewing direction.
Results and Discussions

Cooking softens the structure of initially dry pasta and is normally stopped once a desired texture is reached. If the pasta is not consumed directly, texture continues to change.

Analysing water absorption, water distribution and microstructure during and after cooking revealed that structural, texture-affecting transformations happen independent of the raw materials used. These general transformations during cooking and post-cooking are described first and how they can be monitored combining MRI and light microscopy. Texture changes were analysed in particular during warm-holding of pasta.

Next, the effects of raw materials on these transformations are discussed. They will be discussed separately for the parameters starch and protein content as well as bran addition and bran particle size.

General structural transformations

Cooking: Water absorption and structure transformations

The cooking process is characterised at the macroscale by steady water absorption as shown by weight increase measurement. Water absorption was measured both for industrial-scale and lab-scale spaghetti (data of paper I-III). When cooking time was normalised for the dry spaghetti diameter, water absorption was rather similar for all samples (Figure 6). Comparing the data within a study, differences were even smaller. Water absorption values were recorded at different time points for the various studies, but generally both during early cooking and overcooking stage which started roughly at 4 min×mm\(^{-2}\) (based on determination of optimal cooking times). Lab-scale spaghetti samples varying in protein and starch content (paper II) showed the highest water absorption values at all time points. The higher values might be due to producing reconstituted spaghetti samples by mixing semolina with pure starch or gluten powder. Others have shown that reconstituting starch and gluten to the same amounts as in semolina still alters the pasta structure and results in higher water absorption (Delcour et al., 2000b).

![Figure 6](image-url)
In accordance with these findings, D100 – being produced with unreconstituted durum wheat flour – showed the lowest absorption values within the series of paper II. Except for the spaghetti with the strongly decreased protein content S40D60 showing higher absorption values at all time points, water absorption varied first at the overcooking stage. The existence of a “master curve” implies that the water absorption process during the early cooking stage is foremost governed by the geometry (form and diameter) of the dry pasta product and only to a lesser extent by the raw material. These results are in line with previous research although not all studies linked their results to the diameter (Cafieri et al., 2010; Del Nobile et al., 2005, 2003b; Ogawa et al., 2011; Sekiyama et al., 2012). However, others reported that water uptake differed depending on raw materials (Martinez et al., 2007; Sozer and Dalgic, 2007). Comparing the presented data is complicated as the dry diameter is often not mentioned and only one water absorption value is measured at optimal cooking time. Nevertheless, Rakhesh et al. (2015) showed in controlled studies that soluble fibres such as inulin increase the amount of absorbed water and decrease OCT. The water absorption might also be influenced by process parameters such as drying temperature. Conflicting results have been reported on whether or not water absorption differs with increasing drying temperature (Aimoto et al., 2013; Ogawa and Adachi, 2014).

In a final test, the density of dry spaghetti was estimated (density defined as weight divided by volume based on estimated dry diameter). D100 had a density of about 1.4 g×cm\(^{-3}\), while S40D60, G20D80 and G40D60 had a density of about 1.3 g×cm\(^{-3}\). For comparison, the density for the DS-series was estimated to 1.5 g×cm\(^{-3}\). Thus, density might be another factor for water absorption as lower density of dry spaghetti correlated with higher water absorption values.

To summarise: The raw materials studied in this work did not affect water absorption, but raw materials with highly different hydration properties might do.

The water absorption during cooking is accompanied and likely driven by a steady starch gelatinisation. In fact, Ogawa and Adachi (2013) argue that starch gelatinisation regulates the water ingress into the ungelatinised regions of pasta during cooking. This hypothesis was supported by real-time MRI maps showing that water progressed over time steadily and radially from the surface to the core. A complete time series of the MRI measurement is shown in Figure 6B, and images for selected time points in Figure 7.

**Figure 7** \(T_2\) maps (MRI) and micrographs (PLM) illustrating the progressing water ingress and subsequent starch gelatinisation during cooking and warm-holding. Chart illustrates decrease of ungelatinised, inner core over cooking time as a measure for the water ingress rate. Numbers within the images indicate cooking and warm-holding time, respectively. Images and chart for cooking are from paper II, images for warm-holding from paper I and IV.
An increasing moisture content (as shown by higher $I_0$ values) was accompanied by increasing water-macromolecule interactions (represented by higher $T_2$ values). Corresponding PLM images showed that the area of low $T_2$ values in the spaghetti core correlated with the area of ungelatinised starch (Figure 7). The water ingress as shown by MRI was reflected by progressing starch gelatinisation. A quantitative estimation of the inner, ungelatinised core (MRI and PLM data agreed well) revealed that the core decreased linearly at an early cooking stage (diagram in Figure 7). At a later stage, radius estimation became more uncertain, but still indicated an accelerated decrease of the ungelatinised core. The water ingress was independent on protein and starch content in the sample studied and in line with findings by Del Nobile et al. (2003b) that stated a sharp moisture increase at the gelatinisation front.

**Microstructure of cooked spaghetti**

The correlation between the microstructure as shown by polarized (PLM) and bright-field light microscopy (BFLM) and MRI parameters was analysed in more detail in cooked spaghetti (Figure 8). A micrograph of stained, dry spaghetti is shown for comparison.

Dry spaghetti was characterised by a homogeneous microstructure composed of starch granules embedded in a gluten network. Cooking increased spaghetti in size and created a continuous gradient in the microstructure which can be divided into three regions (core, intermediate and surface region, Heneen and Brismar, 2003).

The microstructure in the core remained unchanged compared to the dry state. Starch granules were not swollen or only very limited (BFLM) and did not gelatinise (PLM). The lower $I_0$ values in the core region (accompanied by very low $T_2*$ values) revealed that the water front had not reached the core thus preventing enough moisture for starch gelatinisation.

![Figure 8](image)

**Figure 8** Cross-sections and representative parts of these cross-sections from core to surface of dry and cooked spaghetti (exemplary shown DS). All light micrographs and MRI maps were derived from separate, individually cooked spaghetti strands. $T_2*$ and $I_0$ maps are compressed to half the size in height to show more voxels. In BFLM, protein appeared green; starch blue/violet; fibre particles were not stained.
In the intermediate region, starch granules increased in size while maintaining their granular form. They lost their birefringence almost completely. Swelling continued, starch granules started to disintegrate and amylose got concentrated at the surface and at the inner core of the granules. In the surface region, starch granules phase separated into amylose and amylopectin. Parts of amylose leached into cooking water, while amylopectin formed random structures.

Fibre particles did not influence these general transformations. Their influence on the microstructure, however, will be discussed later. The gluten network was continuous both in the core and intermediate region and remained intact even in the surface region despite the disintegrated starch granules.

The increasing starch swelling from core to surface was mirrored by gradually increasing $T_2^*$ values. In agreement with previous studies, $T_2^*$ seems to be a good indicator for the degree of starch swelling and gelatinisation (Sekiyama et al., 2012). $I_0$ and thus the moisture content increased only slightly from intermediate to surface region which was consistent with previous results (Del Nobile et al., 2003b).

Furthermore, the radial $I_0$ profiles generally showed only a slight decrease from the surface to the intermediate region. $I_0$ dropped sharply only in the core region (Figure 9). In fact, the radial profiles of $I_0$ values deviated from the profiles of $T_2^*$ for all samples studied (profiles shown for spaghetti of paper I in Figure 9). This illustrated that $T_2^*$ values rather reflected the interaction of water molecules with the local microstructure created of various raw materials. It also proves that $T_2^*$ values did not correlate with water content (represented by $I_0$ values) and confirms that $T_2^*$ should not be used to quantify the water content in cooked spaghetti.

**Post-cooking: Warm-holding**

Starch gelatinisation continued during warm-holding at $80^\circ$C although it happened at a slower pace compared to during cooking. 7 min cooked spaghetti showed a distinct core of ungelatinised starch directly after cooking and still some ungelatinised starch granules after 15 min warm-holding (Figure 7). In comparison to 15 min warm-holding, the ungelatinised core decreased almost to the same extent when cooked for just another 3 min. Concerning starch gelatinisation, cooking and warm-holding are similar processes which differ in the amount of water available. These findings confirm a proposed model of a moving gelatinisation front during warm-holding which was established for rice grains (Briffaz et al., 2014b).

The ongoing starch gelatinisation during cooking and warm-holding affected texture properties, as both processes reduced spaghetti firmness (Table 2). Although a definitive maximum value could not be determined for 7 min cooked spaghetti (probably due to the remaining large
Results and Discussions

25

Figure 10 Force-distance curves to determine firmness (defined as peak force) of warm-held spaghetti (exemplary shown for DS3). Times indicate warm-holding durations. Asterisk (*) marks the incomplete, randomly formed force-top of 7 min cooked spaghetti measured directly after cooking.

un gelatinised core), the initial slope of the force-distance-curve was higher for spaghetti cooked for 7 min compared to 10 min (Figure 10). Independent of cooking time and product, firmness was lower for warm-held spaghetti, but did not differ between 15 min and 90 min holding time (Table 2). Spaghetti at the bottom of the pan had the lowest firmness as it absorbed the surplus water accumulating at the pan bottom (as evidenced by a spaghetti diameter increase), which prolonged the starch gelatinisation resulting in softer spaghetti.

As a side note: The acquired data allowed comparing firmness of 10 min cooked spaghetti that was cooled directly in ice water and spaghetti that was drained for 1 min before cooling (due to the handling of about 2 kg cooked spaghetti). Firmness values decreased with at least 10% within the 1 min (Table 2). The decrease was partly caused by a slightly increased cooking time as it was impossible to empty the cooking pot in a second. It still illustrates that warm-holding of pasta affected texture even in the short term. Operation practices in food services might therefore be more important than the choice of raw material.

Warm-held spaghetti was not only softer, but also stickier than spaghetti directly after cooking (Table 3). While the products differed in stickiness directly after cooking, they did not differ during warm-holding. Again, this could be attributed to the changing water distribution. Stickiness increases as the spaghetti surface gets drier (Dexter et al., 1983a; Guan and Seib, 1994; Horigane et al., 2006). At ambient temperatures stickiness increased with time for about 60 min (Guan and Seib, 1994), but as water redistributed much faster during warm-holding a plateau in stickiness was probably reached within the first 15 min.

Table 2 Firmness at various warm-holding stages (paper IV, measured in N)\(^1\)

<table>
<thead>
<tr>
<th>Time</th>
<th>7 min cooked</th>
<th>10 min cooked</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>DS+SW</td>
<td>DS1</td>
</tr>
<tr>
<td>0 min</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>1 min</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>15 min</td>
<td>8.5±0.7(^b)</td>
<td>8.5±0.5(^b)</td>
</tr>
<tr>
<td>90 min</td>
<td>8.3±0.6(^b)</td>
<td>8.6±0.5(^b)</td>
</tr>
<tr>
<td>90 min bottom</td>
<td>4.7±0.2(^au)</td>
<td>4.8±0.5(^au)</td>
</tr>
</tbody>
</table>

\(^1\) Values followed by same small letter (in same column) or capital letter (in same row) are not significantly different (P < 0.05); Asterisk (*) marks non-determinable values

\(^2\) 0 min; Spaghetti cooled in water directly after end of cooking; 1 min: Spaghetti drained for 1 min then cooled in water
Post-cooking: Storing at ambient temperatures

The results for warm-holding showed that, directly after cooking, moisture distribution was not in equilibrium and moisture redistributed during warm-holding. Moisture redistributes at ambient temperatures as well, but only in the already gelatinised region of the pasta (Horigane et al., 2006). We were able to verify the moisture homogenisation in bran-rich spaghetti by comparing $I_0$ histograms of data acquired over short and long measurement times. The $I_0$ distributions as a measure for the water distribution were narrower (decreasing both high and low intensities of $I_0$ with an unchanged peak maximum) when measured over 92 min (Figure 14E) compared to 16 min (Figure 14D). Also at ambient temperature, moisture homogenisation reduces firmness during storage (Gonzalez et al., 2000). Compared to warm-holding, however, the relative decline was less (comparing results in Table 2 with results by Gonzalez et al., 2000).

### Table 3 Stickiness at various warm-holding stages (paper IV, measured in N)$^1$

<table>
<thead>
<tr>
<th>Time</th>
<th>DS+SW</th>
<th>DS1</th>
<th>DS2</th>
<th>DS3</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 min</td>
<td>2.6±0.3$^{aA}$</td>
<td>2.2±0.3$^{aB}$</td>
<td>1.9±0.1$^{aC}$</td>
<td>1.6±0.1$^{aC}$</td>
</tr>
<tr>
<td>15 min</td>
<td>4.8±0.8$^{aA}$</td>
<td>4.4±0.2$^{aA}$</td>
<td>4.3±0.5$^{aA}$</td>
<td>4.2±0.4$^{aA}$</td>
</tr>
<tr>
<td>90 min</td>
<td>4.2±0.1$^{aA}$</td>
<td>4.5±0.4$^{aA}$</td>
<td>5.1±0.5$^{aA}$</td>
<td>4.3±0.1$^{aB}$</td>
</tr>
<tr>
<td>90 min bottom</td>
<td>2.4±0.2$^{aA}$</td>
<td>2.5±0.2$^{aA}$</td>
<td>2.5±0.1$^{aB}$</td>
<td>1.6±0.2$^{aB}$</td>
</tr>
</tbody>
</table>

$^1$Values followed by same small letter (in same column) or capital letter (in same row) are not significantly different (P < 0.05);

The softer texture of the spaghetti at the pan bottom and the wetted surface of the spaghetti (due to the surplus water) could explain the lower stickiness values compared to spaghetti warm-held in the centre of the pan. The two factors have previously been linked to reduced stickiness (Dexter et al., 1983a; Guan and Seib, 1994).

In summary, access to internal or external moisture regulates the ongoing starch gelatinisation during warm-holding. Ongoing starch gelatinisation and accordingly the decrease of the hard, ungelatinised core is the main factor for the firmness decrease. Optimisation of moisture access can maximise the period of peak texture.
Results and Discussions

Effects of raw materials on texture, microstructure, and water distribution in pasta

The most complete multi-scale analysis has been carried out on four industrial-scale spaghetti samples (2011, paper I). The following analyses have been carried out: Macroscopic water absorption was recorded during cooking and overcooking (Figure 6). Texture parameters (breaking strength and firmness) were determined of dry and cooked spaghetti, respectively. Furthermore, the microstructure was analysed of dry (Figure 11) and cooked spaghetti (Figure 12A-C). The results were linked to MRI. Segments of the acquired 3D MRI data sets are presented as 2D T2* maps visualising the water-macromolecule interactions in axial, longitudinal and tangential direction (Figure 12D). Finally, MRI data is also presented in histograms showing the distribution of $T_2$ and $I_0$ values of the whole measured 3D volume (Figure 14B). Each figure will be described in more detail in the sections to follow.

The industrial-scale samples were produced with the same manufacturing process. The samples showed some common patterns such as a homogeneous microstructure in uncooked state. During cooking, this resulted in a similar water absorption rates and after cooking in a similar change in microstructure and water distribution from core to surface region. Still, raw materials affected the microstructure and water distribution on the micro-scale and these variations were sufficient to affect firmness of the cooked spaghetti.

The four industrial-scale spaghetti were based on the same durum wheat flour, but varied due to further ingredients in composition (starch and protein content, fibre content, type of fibre). To study those three parameter in more detail, lab-scale spaghetti were produced with varied starch and protein content, bran content as well as bran particle size. First, the effects of starch and protein content and thereafter the effects of bran addition and bran particle size on pasta properties will be discussed.

Starch and protein content

Texture properties

Out of the four analysed industrial-scale spaghetti, DS and DS+SW were made of different flour blends that varied mainly in starch and protein content. DS was produced solely with durum wheat flour while DS+SW was produced with a blend of durum and soft wheat flour. Soft wheat pasta has been associated with inferior texture properties, being less firm and more sticky than durum wheat pasta (Dexter et al., 1983b, 1981). This has been attributed to an often lower protein content and weaker gluten properties in soft wheat (Cubadda et al., 2007; Dexter et al., 1981; Marchylo et al., 2004).

In fact, the soft wheat supplemented 2011 DS+SW sample had a lower protein content compared to DS. Subsequently and in agreement with previous studies, the 2011 DS+SW showed, compared to 2011 DS, both a lower breaking strength in dry spaghetti as well as a lower firmness in cooked spaghetti (Table 4). However, the 2014 samples DS+SW and DS did not differ in firmness. This could be explained by DS+SW having higher protein and gluten content as well as higher falling number, indicating improved gluten and starch properties.

Firmness and stickiness were also determined during warm-holding. The 2014 DS+SW spaghetti showed firmness values similar to DS1 at all warm-holding times, whereas the two other durum wheat spaghetti DS2 and DS3 had higher firmness values throughout warm-holding (Table 2). Thus, spaghetti partly made of soft wheat flour did not necessarily have to show inferior cooking quality compared to durum wheat spaghetti. Still, while firmness was comparable, stickiness was higher directly after cooking for DS+SW than for any of the durum wheat spaghetti DS1-3 (Table 3).
**Table 4** Breaking strength of dry and firmness of cooked, industrial-scale spaghetti (measured in N). Paper I: all spaghetti were made of the same batch of DS; paper IV: all spaghetti except DS+SW were made of the same batch of DS. Batches of DS differed between paper I and IV; years indicate time of spaghetti production.

<table>
<thead>
<tr>
<th></th>
<th>Paper I - 2011</th>
<th>Firmness</th>
<th>Firmness</th>
<th>Paper IV - 2014</th>
</tr>
</thead>
<tbody>
<tr>
<td>Breaking strength</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DS</td>
<td>5.3±0.3</td>
<td>13.0±0.7</td>
<td>11.8±0.2</td>
<td></td>
</tr>
<tr>
<td>DS+FB</td>
<td>4.9±0.2</td>
<td>13.2±0.3</td>
<td>10.2±0.4</td>
<td></td>
</tr>
<tr>
<td>DS+WG</td>
<td>4.1±0.4</td>
<td>11.3±0.2</td>
<td>10.0±0.2</td>
<td></td>
</tr>
<tr>
<td>DS+SW</td>
<td>4.5±0.3</td>
<td>10.7±0.4</td>
<td>12.1±0.5</td>
<td></td>
</tr>
</tbody>
</table>

Firmness is generally controlled by cohesiveness of the protein network, stickiness by the composition of the surface region (Cunin et al., 1997). An analysis of microstructure and water distribution can therefore give further explanations for the observed texture differences.

**Microstructure** Micrographs of dry, uncooked spaghetti are shown in Figure 11. The microstructure in all samples was built up of large, oval type-A and small, round type-B starch granules being embedded in a continuous gluten network. Type-A starch granules appeared to be larger in DS+SW than in DS which is plausible as starch granules are on average larger in soft wheat starch compared to durum wheat starch (Pérez and Bertoft, 2010; Soh et al., 2006). DS+SW had a lower protein content than DS, which was also visible as a lower share of the protein phase within the images. The denser gluten network of DS thus might explain the higher breaking strength of DS compared to DS+SW (Table 4).

After cooking, DS and DS+SW showed an overall similar, continuous change in microstructure from core to surface (Figure 12A+B). In both samples starch granules were embedded in continuous gluten network and did not gelatinise in the core. Furthermore, starch granules swelled first in the intermediate region and got disintegrated in the surface region. While the area of ungelatinised starch granules was the same, the depth of the surface region of strongly disintegrated starch granules seemed to be larger in DS+SW (Figure 12C). The protein network seemed to be continuous even in the surface region both for DS and DS+SW, but retained its structure slightly better in DS. Micrographs at higher magnification showed that in DS+SW starch granules were particularly swollen to a larger extent in the intermediate region (Figure 3A in paper I). This might be a result of the lower protein content of DS+SW. Alternatively, it just appeared as stronger starch swelling as the soft-wheat rich DS+SW contained more and on average larger starch granules already in the uncooked state.

**Figure 11** Light micrographs of dried, uncooked, industrial-scale spaghetti.
Whether or not it only appeared as stronger swelling, both cases could explain the lower firmness. Heneen and Brismar (2003) reported that starch granules in soft wheat pasta showed stronger swelling and occupied larger areas in cooked spaghetti. In conjunction with an observed fusion of starch granules in the outer region, the protein network was reported to be less continuous which in turn is often associated with lower firmness (Sissons, 2008). In contrary, a higher share of small granules (achieved through added type-B granules) resulted in higher firmness (Soh et al., 2006).

The lab-scale spaghetti varying in starch and protein content could support the above mentioned results. Common to all cooked samples was again a continuous change in microstructure that progressed with increasing cooking time (Figure 13). At 7.6 min, the starch swelling in the outermost region was rather similar between the samples, but starch swelling decreased towards the core to different extents. Granules were swollen throughout the whole cross-section in the starch-rich S40D60, while the other samples showed a more heterogeneous swelling pattern.

Figure 12 Microstructure of 10 min cooked industrial-scale spaghetti by BFLM (A), PLM (B) and $T_2^*$ parameter maps (D). Magnifications show surface region (C). Data from paper I.
At 13.1 min starch gelatinisation and starch swelling progressed in all samples, but again to different extents. Especially S40D60 showed strongly disintegrated starch granules and areas of amylose enrichments close to the surface. The sample D100 lay in between S40D60 and the gluten-rich samples G20D80 and G40D60, where granule disintegration was less visible and leached amylose remained very close to individual starch granules. The latter samples furthermore showed a continuous protein matrix in the surface region. Hence, the starch and protein content clearly affected the microstructure. A higher protein content created a denser protein network that limited starch swelling.

**MRI data** Common to all spaghetti was a radial gradient in $T_2^*$ values with low $T_2^*$ in the core region (dark areas in axial and longitudinal view, Figure 12D). $T_2^*$ increased gradually from the core to the surface and the highest $T_2^*$ values were especially visible in tangential $T_2^*$ maps. As previously discussed, $T_2^*$ maps agree well with the observed microstructure changes and $T_2^*$ can be seen as an indicator for the degree of starch gelatinisation and swelling.

Both DS and DS+SW showed homogeneous tangential $T_2^*$ maps, but DS+SW exhibited higher $T_2^*$ values than DS which supports the stronger starch swelling in DS+SW observed by light microscopy.

Lab-scale spaghetti varying in starch and protein content were analysed by MRI as well. These samples can be compared more easily in histograms presenting the distribution of the $T_2^*$ values of each voxel in the $T_2^*$ maps (Figure 14A; histogram is based on cooking time of 7.6 min; time series of histograms can be found in Figure 5 in paper II). The $T_2$ distributions of the lab-scale spaghetti can further be compared to the data of the industrial-scale spaghetti (Figure 14B).

It should be noted that the temperature during MRI measurement was at 99°C for lab-scale spaghetti and at 25°C for industrial-scale spaghetti. The higher temperature shifted the $T_2$ distribution towards longer $T_2$ values. Relative differences of $T_2$ distributions were not affected, thus a comparison of the lab-scale with the industrial-scale spaghetti DS+SW and DS is still possible.

The low protein, starch-rich S40D60 showed the longest $T_2$ values and the broadest distribution (Figure 14A). Increasing protein content shortened $T_2$ and narrowed the distribution. Such a behaviour has been reported previously (Kojima et al., 2004). Also DS+SW – having a lower protein content than DS – exhibited the highest $T_2^*$ values and a broader distribution than DS (Figure 14B).
Results and Discussions

Figure 14 Histograms showing the distribution of $T_2^*$ and $I_0$ values of spaghetti cooked close to respective OCT. Duration and temperature of the MRI measurement are indicated together with the image of the acquired data. In (A) $I_0$ values were $T_1$-weighted, disallowing a quantitative comparison between samples and therefore no $I_0$ histogram is presented in (A). Data from paper I-III.

Furthermore, $T_2^*$ values were on average shifted towards higher values in DS+SW. Based on the previously established assumption that $T_2^*$ values correlate with the degree of starch gelatinisation, the $T_2^*$ histograms serve as further evidence that lower protein content in pasta results in stronger starch gelatinisation.

Distributions of $I_0$ as a measure of the local water content were generally less affected by raw material variations, which was plausible as the absolute amount of absorbed water did not differ between the respective samples (as evidenced by water absorption measurements, Figure 6). Still, DS+SW showed a broader distribution compared to DS (Figure 14B). Higher $I_0$ values and thus increased local water contents were found at the surface region. This might partly explain the higher stickiness of DS+SW in comparison to DS. Del Nobile et al. (2005) attributed higher stickiness of low protein spaghetti to higher water concentration at the spaghetti surface.
**Summary**

Spaghetti samples that were processed equally absorbed the same amount of water during early cooking stage – independent of starch and protein content. Also the water transport towards the core and initial starch gelatinisation were not affected. Still, higher protein contents limited starch swelling within the gelatinised region. Lower protein content resulted accordingly in stronger starch swelling that induced lower firmness and higher stickiness. The texture properties of cooked spaghetti are therefore governed by the structural transformations in the already gelatinised region.

**Bran addition and bran particle size**

**Texture properties**

DS+FB and DS+WG contained twice as much dietary fibre as the durum wheat flour based DS. The fibre fraction of DS+FB consisted of particles of elongated shape that were on average smaller than 100 µm, while whole-wheat spaghetti DS+WG included bran particles with a flat shape several hundred µm in diameter.

Bran-rich pasta has often been reported to have an inferior cooking quality compared to semolina-based pasta (Sissons and Fellows, 2014). A softer texture has been explained with a disrupting effect of the bran particles on the gluten matrix (Manthey and Schorno, 2002). In line with this observation is a lower breaking strength of dried, uncooked whole-wheat spaghetti DS+WG compared to DS (Table 4). Accordingly, the cooked DS+WG showed lower firmness than DS both for the 2011 and 2014 samples. The fibre-rich DS+FB behaved differently: The breaking strength of DS+FB was similar to DS and firmness was reduced in the 2014 sample, but not in 2011. Similar results have been shown for dried noodles enriched with wheat bran or wheat fibre (Shiau et al., 2012). Thus, the addition of fibre particles does not necessarily reduce firmness, but might depend on particle size and shape.

To test this hypothesis, spaghetti were produced at lab-scale that included increasing amounts of whole-wheat flour of 0%, 50% and 100% (WW0, WW50 and WW100). Furthermore, spaghetti were produced including bran particles of varying sizes. Based on the particle size distributions, the spaghetti samples could be grouped as having larger particles (median particle size 440 and 370 µm) and having smaller particles (median particle size 160 and 90 µm).

Before discussing the results of texture analysis, it should be noted that studying the effect of processing was not a focus of this thesis, but it was obvious that processing had an influence on texture properties. WW0 and WW50 had similar flour compositions to DS and DS+WG, but were produced on lab- instead of industrial-scale. An extra test showed reduced breaking strengths of about 2 N for the WW-samples compared to 5 and 4 N of DS and DS+WG. Similarly, firmness was greatly reduced (results not shown). The lab-scale extrusion created a less dense structure and the lab-scale drying did likely stabilise the protein network to a lesser extent compared to an industrial-scale dryer. Hence, the weakened gluten-starch matrix might hide possible differences caused by bran particle size.

Still, texture analysis revealed that smaller bran particles were associated with a higher breaking strength in dried spaghetti (Table 5). Firmness was also slightly higher for smaller bran particles while stickiness was not affected. Shiau et al. (2012) observed higher breaking strength and firmness for fine bran in noodles compared to coarse bran and attributed it to smaller particles having less adverse effect on gluten network formation. Although studying only a small particle size range, Niu et al. (2014) found increasing hardness with decreasing bran particle size in cooked noodles.
Results and Discussions

Table 5 Breaking strength of dry, firmness and stickiness of 8 min cooked bran-rich spaghetti varying in bran particle size (paper III, measured in N)

<table>
<thead>
<tr>
<th>Sample</th>
<th>Breaking strength</th>
<th>Firmness</th>
<th>Stickiness</th>
</tr>
</thead>
<tbody>
<tr>
<td>S440</td>
<td>1.5±0.1</td>
<td>6.3±0.4</td>
<td>0.61±0.04</td>
</tr>
<tr>
<td>S370</td>
<td>1.8±0.2</td>
<td>6.6±0.2</td>
<td>0.63±0.05</td>
</tr>
<tr>
<td>S160</td>
<td>2.2±0.2</td>
<td>7.0±0.3</td>
<td>0.63±0.06</td>
</tr>
<tr>
<td>S90</td>
<td>2.2±0.2</td>
<td>6.9±0.3</td>
<td>0.60±0.02</td>
</tr>
</tbody>
</table>

A subsequent sensory analysis found almost no effect of bran particle size on elasticity, firmness and stickiness (Table 6). The slight differences in firmness as determined by texture analysis were not large enough to be detected by the panel. Others reported similarly higher firmness with smaller bran particle sizes during texture analysis while sensory analysis did not show significant differences (Chen et al., 2011).

Sensorial observed firmness might not be affected by bran particle size, but overall liking was. The samples S90 and S160 showed higher overall liking, which correlated with lower values for observed surface roughness as well as whole-wheat aroma. Again, these findings agreed with Chen et al. (2011) reporting that smaller particles produced a smoother surface and improved appearance and taste in noodles.

Table 6 Sensory properties of 10 min cooked bran-rich spaghetti varying in bran particle size (paper III); sensory attributes determined as difference from control (scale -5 to +5, control WW50: 0); overall liking without reference (0 to 10)\(^1\)

<table>
<thead>
<tr>
<th>Sample</th>
<th>Elasticity</th>
<th>Firmness</th>
<th>Stickiness</th>
<th>Surface Roughness</th>
<th>Whole-wheat aroma</th>
<th>Overall liking</th>
</tr>
</thead>
<tbody>
<tr>
<td>S440</td>
<td>-1.5(^a)</td>
<td>-1.0(^a)</td>
<td>-0.3(^a)</td>
<td>1.3(^a)</td>
<td>1.1(^a)</td>
<td>3.2(^a)</td>
</tr>
<tr>
<td>S370</td>
<td>-1.1(^ab)</td>
<td>-0.7(^a)</td>
<td>-0.7(^a)</td>
<td>0.8(^a)</td>
<td>0.8(^ab)</td>
<td>4.6(^ab)</td>
</tr>
<tr>
<td>S160</td>
<td>-0.1(^b)</td>
<td>-0.1(^a)</td>
<td>-0.9(^a)</td>
<td>-1.0(^b)</td>
<td>-0.6(^c)</td>
<td>5.4(^d)</td>
</tr>
<tr>
<td>S90</td>
<td>-1.6(^a)</td>
<td>-1.0(^a)</td>
<td>-0.5(^a)</td>
<td>-0.8(^b)</td>
<td>-0.3(^bc)</td>
<td>5.7(^d)</td>
</tr>
</tbody>
</table>

\(^1\)Values followed by same letter in a column are not statistically significant different

Microstructure Generally, bran and fibre particles are visible both in bright-field and polarised light microscopy. The higher dietary fibre content of DS+FB and DS+WG compared to DS and DS+SW was reflected by more visible particles in dried, uncooked spaghetti (Figure 11) and was also observed in cooked spaghetti (Figure 12 A+B). Wheat flour contains minor amounts of bran as well as cell wall material from the endosperm and thus some fibre particles were visible in DS and DS+SW as well. The correlation between the amount of fibre and amount of visible fibre particles was easily observed in the micrographs of the lab-scale spaghetti of varying whole-wheat content (Figure 15A).

![Figure 15](image_url) Polarised micrographs (A) and $T_2^*$ maps (B) of cooked spaghetti varying in whole-wheat content (0, 50 and 100% whole-wheat flour).
Generally, fibre particles got aligned with the extrusion direction due to high shear forces during extrusion. The alignment can be seen in the longitudinal sections as the fibre particles were mostly aligned parallel to each other (Figure 12 A+B). Furthermore, the flat-shaped bran particles appeared in cross sections as bent and folded structures. As the folding happens during extrusion, folded bran particles were visible already in the uncooked spaghetti (large particle to the top right in DS+WG image in Figure 11).

Manthey and Schorno (2002) based their argument of the disrupting effect of bran on pasta matrix on the observation of a gap between a bran particle and the surrounding gluten-starch matrix. In the micrographs of dried, uncooked spaghetti no such gaps were observed, independent whether fibre particles were small or large. The particles were well embedded in the protein-starch matrix (Figure 11). In lab-scale spaghetti, however, some large bran particles detached from the surrounding matrix during cryostat sectioning, while small particles did not. If gaps occur between bran and the protein-starch matrix might therefore depend on the production process.

Maybe “disruption” is a too strong word, but fibre particles affected the gluten network as fibre particles occupied potential spaces for protein in the gluten network (Pavlovich-Abril et al., 2012). This effect is also referred to as diluting the gluten network (Shiau et al., 2012).

Larger bran particles were distinctly distributed and created a heterogeneous microstructure (Figure 11), which might explain the lower breaking strength. DS+FB showed an unchanged breaking strength compared to DS. The small particles of DS+FB were homogeneously distributed throughout the entire pasta matrix. The size of the fibre particles in the cross section was in the same magnitude as large starch granules, and presumably the gluten network was diluted to lesser extent.

In cooked spaghetti, the overall radial change in microstructure was not affected by the presence of fibre particles. The area of ungelatinised starch was the same for DS+WG, DS+FB and DS (Figure 12B), and also starch granules swelled and disintegrated to the same extent in the surface region (Figure 12C). Still, due to the alignment of the fibre particles, the particles were often orientated perpendicular to the direction of the water ingress. Especially large bran particles exhibited a barrier effect as the degree of starch swelling differed in front and behind the bran particle. This behaviour will be described in detail for the lab-scale spaghetti varying in bran particle size.

The lab-scale spaghetti varying in bran-particle size (smaller bran particles S90 and S160, larger bran particles S370 and S440) were analysed in their cooked state (Figure 16). Polarized light micrographs confirmed the correlation between smaller particles and more homogeneous distribution of bran particles throughout the pasta matrix (Figure 16A). Again, mainly large bran particles appeared folded and bent, while smaller particles formed rather straight strings.

Similar to the industrial-scale whole-wheat spaghetti, bran particle size did not affect the commonly observed change in microstructure from core to surface (Figure 16B). Bran particles of all sizes were orientated along the extrusion direction (see longitudinal and tangential sections). Some of the large bran particles were further orientated perpendicular to the radius and were especially visible in the tangential view where the particles formed large surfaces (Figure 16B).

To visualise the orientation of bran particles in detail as well as the orientation effect on local degree of starch swelling, selected magnifications of large bran particles are shown in all three directions (Figure 17). Three bran particles are shown in cross-section (Figure 17a-c). Common to all shown bran particles was that starch granules close to the bran particle and facing towards the spaghetti surface were swollen to a much larger extent than granules facing towards the core (arrows are shown for comparison). The bran particles hindered the water transport towards the core and therefore limited the degree of starch swelling.
Figure 16 PLM (A), BFLM (B) and $T_2^*$ maps (C) of 8 min cooked spaghetti of varying bran particle size. Tangential micrographs (B) and $T_2^*$ maps (C) 200-300 µm away from the spaghetti surface. Asterisks (*) in (B) show artefacts from sample preparation and dark areas are folded sections. Images in (A, B, C) were derived from individually prepared spaghetti strands. Data from paper III.
For comparison: The continuously increasing starch swelling towards the core was observed atop and beneath the bran particles in Figure 17B and C, whereas at the level of the particles the degree of starch swelling changed more abruptly. The local barrier effect on starch swelling was mostly observed near the spaghetti surface, but was also visible further towards the core (see magnifications in Figure 17 A).

Another example for the barrier effect of bran is shown in a longitudinal section (Figure 17I). Median bran particle size of S440 was 440 µm, but some particles were much longer as illustrated in Figure 17K. The bran particle in Figure 17I is shown to illustrate that not all particles were orientated perpendicular towards the radius of the spaghetti. This particle was instead orientated parallel to the radius and affected the water transport only to a very limited degree.

Finally, perpendicular orientated bran particles were easily observed in tangential sections (Figure 17E-H). Also in tangential sections, the barrier effect was evident. The slice shown in Figure 17H1 was located about 30 µm further towards the spaghetti core than slice shown in H2. The beginnings of the large bran particle in H1 can already be seen in H2. The starch granules atop the bran particle were swollen to a larger extent than starch granules alongside the particle (see arrows for comparison).

The particles were often even larger than visualised by the images as a 5 µm thick slice cannot reproduce the true 3D structure of a particle. The bran particles in the centre and in the lower left corner of Figure 17G were in fact belonging to the very same particle, which formed a surface of at least 0.5mm×1mm. The planar shape of the particles as well as the thin microscopy slices allowed to show single bran layers such as likely tube cells (Figure 17D), cells belonging to the pericarp (Figure 17F) as well as aleurone layer combined with nucellar layer (Figure 17G). The layers were attributed according to Slavin et al. (2000) and Surget and Barron (2005). Previous research did not focus on the orientation of bran particles, but at least one study also showed aleurone cells (similar to Figure 17G) at the pasta surface (Manthey and Schorno, 2002).

Figure 17 Bran particles in cooked spaghetti in cross section (a-c), tangential (e-h) and longitudinal sections (i-k). a2 and a3 show magnified areas of a1. Slices in h1 and h2 are about 30 µm apart. Arrows indicate different degree of starch swelling. Data from paper I and III.
**MRI data**

All samples, both industrial-scale and lab-scale spaghetti, showed $T_2^*$ maps where $T_2^*$ values increased from the core to the surface (Figure 12D, Figure 15B, Figure 16C). This reflects that bran addition did not affect the overall changes in microstructure in cooked spaghetti.

The heterogeneous microstructure of the whole-wheat spaghetti DS+WG, containing large bran particles, was reflected in the $T_2^*$ maps (Figure 12D). DS+WG showed overall lowered $T_2^*$ values in all directions compared to DS and also DS+FB, but the heterogeneity became most evident in the tangential view. Large areas of low $T_2^*$ values were surround by areas of higher $T_2^*$ values. As these dark areas were also elongated and orientated along the extrusion direction, we assumed that these areas represented the bran particles. The reduced $T_2^*$ could be due to faster exchange of water protons with exchangeable fibre protons or due to the less swelled starch granules/low water content close to the fibre particles or a mix of both reasons. Interestingly, $T_2^*$ maps of DS and DS+FB were similar, although DS+FB contained twice as much fibre as DS. This could indicate that the small fibre particles of DS+FB interacted less with water protons compared to starch and gluten and they did not affect starch swelling locally. Alternatively, small fibre particles are distributed so homogeneously throughout the pasta matrix that their effect on $T_2^*$ could not be captured with the voxel size of 78µm$^3$ of the MRI measurement. The $T_2^*$ value in each voxel is an average of all components such as starch, gluten and fibre within the voxel volume. In the case of DS+FB, there might be too few fibre particles in each voxel volume to affect $T_2^*$.

The overall reduced $T_2^*$ values can be explained by the addition of bran particles. $T_2^*$ values shifted towards shorter values over the whole radial profile in the $T_2^*$ maps shown for increasing whole-wheat content (Figure 15B).

Heterogeneity in $T_2^*$ maps can be connected to bran particle size. Similar to DS+WG, the large bran particles containing S440 and S370 showed more heterogeneous $T_2^*$ maps than S160 and S90. Again, the largest bran particle size resulted in the most heterogeneous tangential $T_2^*$ map.

Industrial-scale spaghetti and lab-scale spaghetti also showed same trends when analysing the distributions of $T_2^*$ and $I_0$ (Figure 14B-D). Increasing whole-wheat content (WW0, WW50, WW100) shifted the $T_2^*$ distribution towards shorter $T_2^*$ values and decreased especially the amount of high $T_2^*$ values (Figure 14C). The bran-rich DS+WG similarly showed a $T_2^*$ distribution shifted towards lower $T_2^*$ values compared to DS and DS+FB, which exhibited a similar pattern of $T_2^*$ values (Figure 14B).

Comparable lowering of $T_2^*$ values has been reported for increasing whole-wheat contents in dough systems and has been attributed to a moisture migration from the gluten network to the bran particles (Bock et al., 2013; Li et al., 2014). Accordingly, Chen et al. (2011) speculated that wheat bran absorbs water during the cooking process of noodles and thereby influences water transport to starch granules near to bran and hence the swelling behaviour of that starch. However, it is questionable whether bran particles absorb water from the surrounding gluten-starch matrix during cooking. $I_0$ distributions were broadened for increasing whole-wheat content (Figure 14C). $I_0$ being a measure for the local water content, this meant that water was distributed more heterogeneously in spaghetti with high bran content. Our interpretation is that bran particles did not absorb water during cooking or at least not to the same extent as the surrounding gluten-starch matrix. Instead, large bran particles redistributed the water around the particles.

Bran particle size did not affect $T_2^*$ maximum peak in lab-scale spaghetti, though larger bran particles broadened the $T_2^*$ distribution (Figure 14D). A broader $T_2^*$ distribution indicates a more heterogeneous microstructure and might also be related to a more uneven degree of starch gelatinisation. Again, a similar behaviour was observed when comparing DS, DS+FB...
Effects of: Bran addition and bran particle size

and DS+WG. The bran-rich DS+WG had the broadest $T_2^*$ distribution, while DS+FB (enriched with smaller fibre particles) had a slightly broader distribution than the control DS (Figure 14B). $I_0$ distributions were affected less by bran particle size, but larger bran particles in lab-scale and industrial-scale spaghetti still showed slightly broadened $I_0$ distributions (Figure 14B+C). This supports the results of increasing whole-wheat content, such as that large bran particles redistributed water to a stronger degree than smaller bran particles due to their size.

Summary Texture and sensory properties of whole-wheat spaghetti can be modulated by bran particle size. Microscopy and MRI gave insights to explain the observed variations in texture. The macrostructure of cooked spaghetti was not affected by bran particle sizes, but larger bran particles induced a more heterogeneous microstructure. During cooking, bran particles absorbed water to a lesser extent than the surrounding starch-gluten matrix. Large particles exhibited further a barrier effect for water migration and thereby locally limited the degree of starch gelatinisation.
Conclusions

The main objective of this work was to characterise the interplay of water and microstructure in pasta as affected by choice of raw materials.

To do so, magnetic resonance imaging was adapted to map the water distribution in 3D at high resolution. Moreover, MRI and light microscopy complemented each other. Thereby, variations in water distribution were linked to structural components such as starch, gluten and fibre. Analysing the microstructure from different perspectives was demonstrated to be essential in visualising the orientation of anisotropic structures such as fibre particles. A further adapted MRI method allowed monitoring the cooking process of pasta in real-time.

The cooking process was characterised by steady water absorption and the water ingress towards the ungelatinised core was regulated by starch gelatinisation. It was shown that macroscopic water absorption as well as water ingress was governed by pasta geometry and depended only to a limited degree on raw materials. However, the gelatinised region was affected by raw material composition as an increased protein content limited starch swelling.

The influence of fibre particles was characterized by studying bran particles of varying particle size. Bran particles get aligned during extrusion, which often results in a perpendicular orientation compared to the direction of the water ingress. Thereby, bran particles can create barriers for the water ingress and redistributes the water around the particles. Connected to the redistribution is the local effect of restricting starch swelling. The severity of the barrier effect increases with particle size. Larger bran particles alter the microstructure already in dry pasta. Both factors can partly explain why, in sensory analysis, smaller bran particles were preferred as they alter the microstructure to a lesser extent.

After cooking, water distribution was not in equilibrium within pasta. Instead, the present internal moisture was sufficient to continue starch gelatinisation during warm-holding. This resulted in a fast decrease in firmness and increase in stickiness. When the moisture distribution reached its equilibrium, texture properties did not change with longer warm-holding times. These findings show that cooking and warm-holding are similar processes that only differ in the amount of water available.
**Future research**

Working with complex food structures such as pasta allows for various research approaches. The research in this work touched different areas and it is therefore of interest both to focus on further method development as well as a more detailed analysis of raw materials and their processing.

The described interplay of water and microstructure was based on correlations as the techniques were applied to different spaghetti strands of the same batch. It would be interesting to carry out both MRI and light microscopy on the very same spaghetti strand. This has not been tested in practical terms, but it is theoretically imaginable to freeze the spaghetti strand immediately after the MRI measurement. The spaghetti strand could then be cryo-sectioned throughout the sample, which could serve as a base for a 3D reconstruction based on light microscopy. Comparing 3D data obtained by MRI and light microscopy from the very same spaghetti strand could give more complete understanding of the interdependence of water distribution and microstructure.

We were able to study water migration during cooking by MRI in real-time and the same method can in principle be applied to study the water migration during warm-holding as well. However, there are some practical challenges to overcome.

Another approach could be to combine MRI and light microscopy with further spatially resolved methods such as fluorescence fingerprint imaging (Kokawa et al., 2014), microtomography (Zhang et al., 2013) as well as moisture maps based on photography and image processing (Ogawa and Adachi, 2013). Confocal laser scanning microscopy is particularly suitable to visualise the protein network in pasta (Zweifel et al., 2003). It would be interesting to use it with quantum dots as fluorescent probe enabling to study the protein network in more clarity (Sozer and Kokini, 2014). Even non-spatially resolved methods can give further valuable information such as differential scanning calorimetry that can be used to determine the degree of starch gelatinisation within a pasta sample.

The described methods can also be used to study how raw materials affect and are affected by the production process in more detail. Especially for bran and fibre particles it could be studied if and how they attract water during mixing and drying.

In keeping with numerous studies, the new raw material – in our case bran – should not interfere with the protein network to achieve the best texture. Milling bran to smaller particle size was only one option to address this challenge. To achieve a pronounced increase in quality of whole-wheat pasta, solely optimising the milling of bran might not be enough. Instead, a combination with other processes should be considered. Several processes have been suggested recently and it would be interesting to see them applied in a pasta context. These processes included microfluidized (Mert et al., 2014), tribocharged (Chen et al., 2014) and cryogenic ground bran (Hemery et al., 2010), as well as heat treated (Sudha et al., 2011) or fermented bran (Coda et al., 2014). To increase complexity even more: While optimising the processing of bran for improved pasta texture properties, the potential nutritional benefits of bran may not be compromised.

Finally, the research on warm-held pasta revealed that the texture deterioration is caused by a surplus of water – both inside and outside of the pasta. Besides finding raw materials that tolerate warm-holding, it is advisable to adapt the post-cooking handling of pasta so that excess water is removed as fast as possible.
Acknowledgements

I would like to thank everyone, both colleagues, friends and family, who have supported me during my time as a PhD student.

First of all, a very special thank you to my supervisor Maud Langton for your encouragement, support and understanding throughout the years.

Thank you, Annelie Moldin, for taking the time to be a co-supervisor and for all good pasta discussions, for your help and for introducing me to the secrets of pasta making.

Diana Bernin: It is fair to say that, without you, this work would not look the same. How would the title figure look like without the left part? Thank you for your patience in teaching me MRI, your sincere interest in pasta research, your curiosity, commitment and endurance. Where do you find all your energy?

I want to thank Lantmännen Research Foundation for initiating this research project. Special thanks to Carina Bohman, Christian Malmberg, Sofie Villman and Ingmar Börjesson for your support, your ideas and all the fruitful discussions. I would like to further acknowledge the practical help of the many Lantmännen employees in Malmö, Järna, Stockholm and Uppsala.

A special thanks to my mentor Mats Larsson for interesting discussions and for giving me insights into the wider implications of research for the food industry.

I thank SuMo Biomaterials for a stimulating research environment as well as for interesting meetings and courses within the Centre. Special thanks to Magnus Nydén for introducing me to Diana and suggesting MRI “might be something to try one day”.

Thank you for good collaboration to Magnus Röding and Daniel Topgaard.

It was a pleasure to work with my master thesis student Emmy Sandberg.

I am grateful to Camilla Öhgren and Niklas Lorén for reviewing this work and valuable comments.

Thank you to all kind colleagues at SP Food and Bioscience (formerly known as SIK) and especially to the folks at the Structure and Material design group. You made me always feel at home and did not forget me even after I partly found another office room in Uppsala at the Department of Food Science, Swedish University of Agricultural Sciences (SLU). My regards go to all colleagues at the department, but especially to the superb Structure and Properties group.

A special thanks to Daniel Johansson, my roommate from coast to coast.

I would like to thank all PhD students at SIK and SLU for good company and enjoyable study trips, together with the Research Schools LiFT and Food in Focus.

Finally, a deep thank you to one more PhD student who supported me the most: Babett, you are awesome!
References


