

Biotest Group: Creating Value. Living Values



German Healthcare Conference, Zürich Biotest AG

September 8, 2010



Disclaimer

This document contains forward-looking statements on overall economic development as well as on the business, earnings, financial and asset situation of Biotest AG and its subsidiaries. These statements are based on current plans, estimates, forecasts and expectations of the company and thus are subject to risks and elements of uncertainty that could result in deviation of actual developments from expected developments.

The forward-looking statements are only valid at the time of publication. Biotest does not intend to update the forward-looking statements and assumes no obligation to do so.

All figures reported relate to the Continuing Operations of the Biotest Group after the disposal of the transfusion and transplantation diagnostic activities to Bio-Rad Laboratories Inc. These activities are being reported as Discontinued Operations. With the exception of the statement of financial position, the previous year's figures have been adjusted accordingly.

All comparative figures relate to the corresponding last year's period, unless stated otherwise.



Biotest at a glance

Key Figures: FY 2009 H1 2010

 Sales
 € 438.6 m (+14.2%)
 € 227.1 m (+4.0%)

 Thereof Plasma Proteins
 € 390.1 m (+14.9%)
 € 200.6 m (+3.2%)

 EBIT
 € 61.6 m (+4.6%)
 € 23.7 m (-24.0%)

Business sectors

Pharmaceuticals

Diagnostics

Divisions

Plasma Proteins

- Immunoglobulins
- Hyper-immunoglobulins
- Clotting factors
- Albumin

Biotherapeutics

 Monoclonal antibodies

Microbiological Monitoring

Hygiene monitoring





Financials H1 2010



H1 2010 – At a glance

- H1 Sales increase + 4.0% to €227.1 million in difficult market environment.
- Continued influences on EBIT:
 - further price decrease for plasma protein products
 - continued unabsorbed costs in US (finalisation production facility Boca Raton)
 - increased R&D expenses: €4.1 million (+19%) incl. consistency batches at BPC and regulatory filing for BLA BivigamTM
- H1 EBIT € 23.7 million (-24%)
- Revised EBIT Outlook











Expectations FY 2010

- Sales growth in lower single digit range
- Further price pressures expected for Intratect and Haemoctin
- Negative impact by German Healthcare Reform
- Continued unabsorbed costs in US (production facility Boca Raton)
- Shifting of products in higher margin markets not successful
 - EBIT level of 2009 will not be reached



New EBIT guidance: €45 million +/- 10%

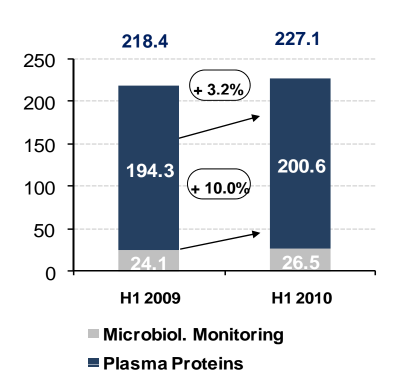
EBIT Guidance incl. Discontinued Operations:

€45 million +/- 10% plus €18 million



Sales growth despite difficult environment

Sales of Plasma Proteins & Microbiological Monitoring (€m)



- Sales in the first half year of 2010 were up by 4.0% to 227.1 million vs. H1 2009
- The Microbiological Monitoring segment increased by a rate of 10.0 %, mainly through products manufactured by heipha
- The Group's Plasma Proteins business grew with 3.2%
- Robust performance in challenging business environment



Sales Plasma Proteins

| Sales Plasma Proteins H1 2009 | € | 194.3 m |
|-------------------------------|---|---------|
| Volume effect | € | 20.9 m |
| Price effect | € | -14.6 m |
| | | |
| Sales Plasma Proteins H1 2010 | | 200 C |
| Sales Plasma Proteins HI 2010 | € | 200.6 m |

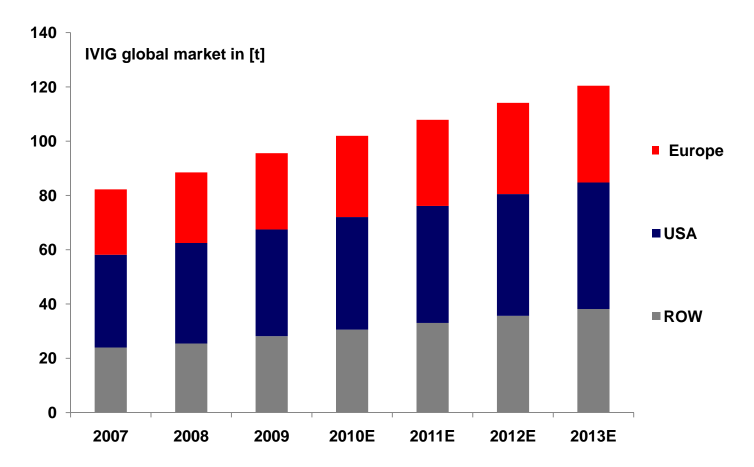




Plasma Proteins



Development of IVIG markets by regions



• The IVIG market will continue to grow (5% p.a.), particularly by increased demand in emerging markets

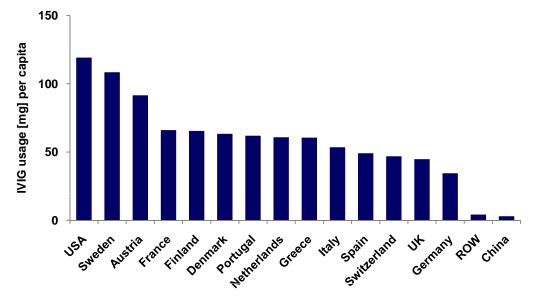
Source: MRB, Analyst Reports, Biotest Market Research



Further growth of immunoglobulin market expected

Demand growth driven by

- Favorable demographics: age, weight
- Improved diagnosis, higher dosing level and longer time on therapy
- Continued clinical evidence supporting established and new indications
- Geographical expansion



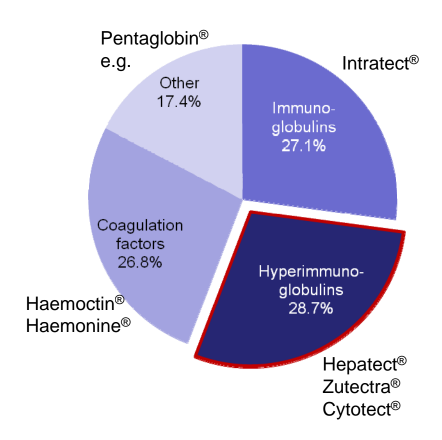
Source: Global Insight, MRB, PPTA, APFA

^{*:} Poster: "A European, multicentre, open and prospective study on clinical efficacy, safety, and pharmacological properties of Intratect® (human normal immunoglobulin for iv administration) in patients with primary immunodeficiency (PID)"; E. Bernatowska et al., 2006



Biotest: A market leader in special preparations

Biotest plasma proteins in 2009: sales by product category



Hyperimmunoglobulins and special preparations are a very attractive segment:

- Stable prices
- High market entry barriers
- Biotest is totally self-sufficient in hyperimmune plasma procurement









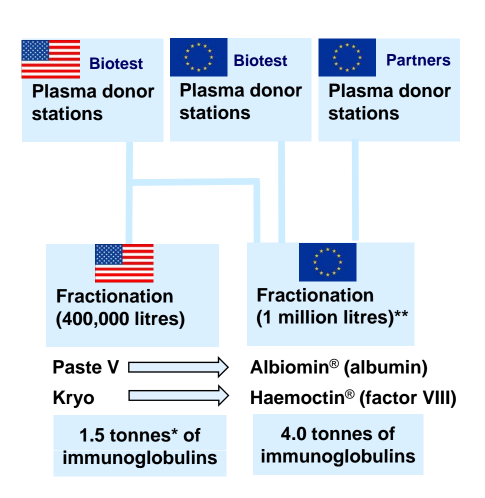
US manufacturing plant

- State-of-the-art manufacturing facility at Biotest Pharmaceuticals Corp. (BPC) in Boca Raton, Florida
- Fractionation: 400,000 litres per annum
- Immunoglobulin production: 1.5 tonnes per annum
- Plasma collection at 11 BPC-owned plasma collection centres





Plasma Proteins – Efficient production network



- 21 plasma collection centres
- Level of self-sufficiency: 40% for standard plasma
- Exchange of intermediate products from US to Europe from end of 2010
- Network increases EBIT margin

^{*} Approval will probably be granted end of 2011

^{**} Production in Dreieich and capacities at partners



Major R&D progress of Plasma Protein projects



Zutectra®

Post approval trial to examine convenience and self-medication at home with 70 patients



IgM Concentrate

Phase II to start mid of 2011 Treatment of serious bacterial infections High functional activity, good tolerability



Cytotect®:

Phase III clinical trial ongoing Prevention of prenatal CMV infection Interim evaluation planned for end of 2010

Hepatitis B immunglobulin (subcutaneous / intramuscular) in neonates

Phase III trial, recruitment completed Final Draft of Study Report end of 2010 Marketing approval: aiming at Germany first



From Nature for Life



Biotherapeutics



Biotherapeutics: Focused research

Biotherapeutics: Focused research

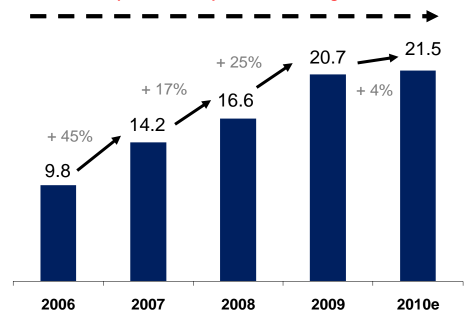
- High medical need
- Rapidly growing markets
- Blockbuster potential

R&D expense – Biotherapeutics (in €million)

Cap on Biotherapeutics R&D budget

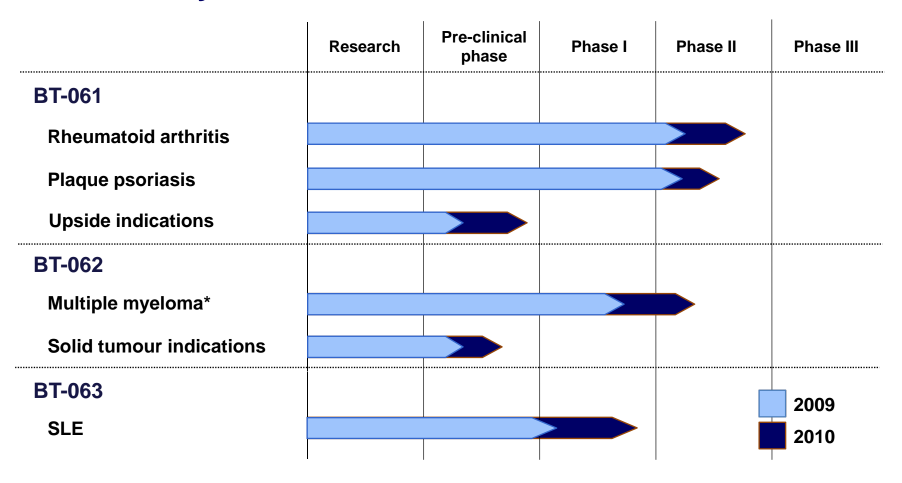
Lead indications

| BT-061 | Rheumatoid Arthritis, Psoriasis | | | | |
|--------|------------------------------------|--|--|--|--|
| BT-062 | Multiple Myeloma | | | | |
| BT-063 | Systemic Lupus Erythematosus | | | | |





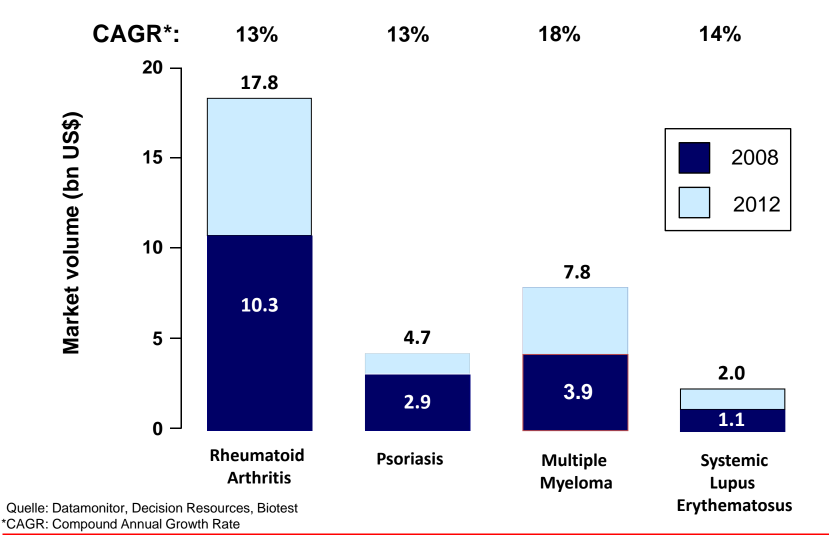
Biotherapeutics: Significant project progress in financial year 2009 and 2010



^{*} Phase I/IIa clinical trial approved by FDA (IND)



Biotherapeutics: Continuously growing market potential





Rheumatoid Arthritis: Competitive market environment Favourable positioning is key to success

| | Cytokine neutralizing (TNF α and others) | Targeting B cells or T cells | Targeting Tregs: BT-061 |
|--------------------------|---|---|---|
| MoA ¹⁾ | Neutralization of cytokines | Depletion/inactivation of immune cells | Selective activation of Tregs |
| Weakness/ Threats | Black box warning: risk of infection and malignancy FDA alert for: invasive fungal infections and increased risk of lymphoma in children | Black box warning for PML²⁾ Increased risk of infection B-cell depletion (up to 1 yr) Severe infusion reactions | • Late market entry requires clear USP ³⁾ and positioning |
| Strength/ Opportunity | Market dominanceBroad safety database | Treatment of TNF non responders | Superior efficacy expected Mode of action supports good safety profile (no signs of immunosuppression, cytokine release or lymphocyte depletion) |

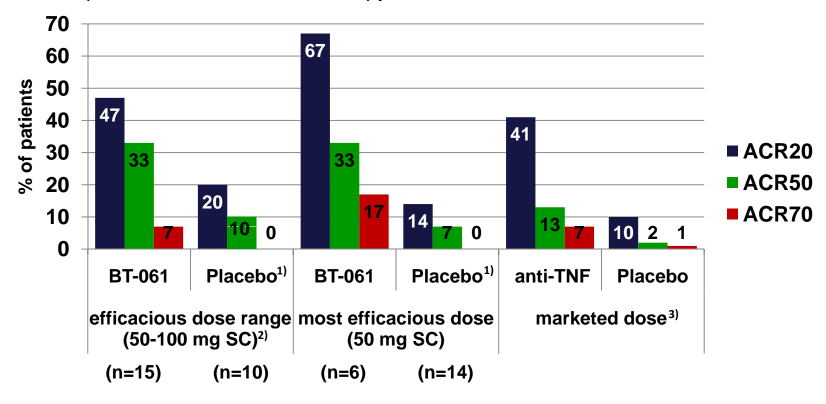
Positioning of BT-061 by new MoA, which translates into superior efficacy and safety

¹⁾ Mode of Action ²⁾ Progressive multifocal leucoencephalopathy ³⁾ Unique selling point



Repeated treatment of RA patients with BT-061 (monotherapy) Benchmarking against gold standard of biologic therapy

ACR responses at week 7, monotherapy



¹⁾ Two patients from each completed SC dose group; 2) Only patients that received all treatments over the 6 week periode

³⁾ Phase III trial results of anti-TNF monotherapy in DMARD non-responders at week 7



Biotherapeutics: Established own production capacities





Development structures in the segment:

- GMP production of monoclonal antibodies established in Boca Raton (BPC)
- Manufactured first large-scale batches of BT-061 in own production facility
- Start of GMP production of BT-062 at BPC in 2011



BT-061 partnership



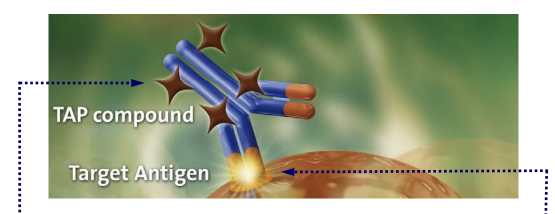
Biotest strategy:

Cooperation with partner from clinical phase III

- Negotiations with international pharmaceutical companies ongoing
- High level of interest
- Request for confirmation of positive trial results via further phase II clinical trials
- Stand-alone further development of mAb until agreement is reached



Competitive edge BT-062: Intrinsic properties provide basis for product positioning



Toxin moiety mediates high efficacy

- High potency independent of patient's immune system
- Toxin technology with best track record: Sanofi Aventis, Biogen Idec, Bayer, Roche/Genentech amongst licensees
- First filing of TAP¹⁾ mAb expected in 2010 (Genentech)

Antibody moiety mediates high specificity

- Unique targeting to CD138
- CD138 highly overexpressed in MM and other cancer cells
- CD138 not expressed on bone marrow stroma cells
- Good tolerability up to 160 mg/m²

1) TAP: Tumor activated payload



BT-062: Single-dose study 969 in Multiple Myeloma First efficacy data, August 2010

| Number of patients | Total | Percentage | Objective response | Clinical benefit (%) |
|--------------------------------------|-------|------------|--------------------|-------------------------|
| treated with BT-062* | 32 | | | |
| efficacy data available | 25 | 100% | | |
| - disease progression within 6 weeks | 11 | 44% | | |
| - stable disease ≥ 9 weeks | 12 | 48% | | |
| - minor response | 1 | 4% | 00/ | 56% |
| - partial response | 1 | 4% | 8% | |

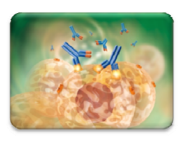
- > BT-062 shows anti-tumor activity already in repeated single dose schedule
- > Further patients were enrolled in MTD** cohort up to a total of 13

^{*}Median number of prior chemotherapies: 7 (range: 2-15); 33% of patients had 10 or more prior chemotherapies

^{**}MTD: Maximum tolerated dose; Response criteria as defined by International Myeloma Working Group

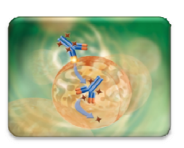


Outlook Biotherapeutics: Next steps in clinical development initiated



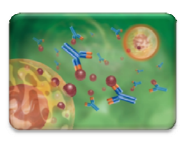
BT-061:

- First encouraging clinical data from both lead indications
- Phase II trial in Psoriasis started
- Phase IIb in RA initiated
- Discussion with strategic partners ongoing



BT-062:

- First indications of efficacy from dose-escalating study
- Multiple dose phase I/IIa trial approved by FDA
- Study initiated



BT-063:

- Phase I study approved in Sept. 2009
- Treatment at 7th dose level completed (02 2010)





Outlook for 2010



Further outlook Biotest Group

- Despite difficult business environment we continue to invest into R&D of Plasma Protein Projects and Biotherapeutics
- Full pipeline of Plasma Protein products and Biotherapeutics with a potential to reach the market within the next years
- BPC/ USA: access to the single biggest plasma protein market
 - ➤ Q3 2010 BLA submission of BivigamTM on track
 - ➤ Launch of BivigamTM (IVIG) expected to take place in H2 2011
 - Additional market potential of \$ 100 million



Contact and Financial Calendar 2010/2011

Investor Relations Biotest AG:

Dr. Monika Buttkereit
Head of Investor Relations

Phone: +49 (0) 6103 - 801 -4406 Fax: +49 (0) 6103 - 801 -347

E-Mail: investor_relations@biotest.de

Financial Calendar 2010/ 2011 Nov 08, 2010 **Q3 Report 2010/ Analyst's Conference** Mar 22, 2011 FY 2010/ **Analyst conference** May 10, 2011 **Q1 Report 2011** May 12, 2011 **Annual General Meeting** Aug 11, 2011 **Q2 Report 2011** Nov 10, 2011 **Q3 Report 2011/ Analyst conference**