Improved Immunohistochemical Detection of Type 1 Insulin-Like Growth Factor Receptor in Human Tumors

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Abstract

Background: Insulin-like growth factors (IGFs) are known to play important roles in cancer biology, prompting evaluation of drugs targeting type 1 IGF receptor (IGF-1R). However, there is considerable lack of consensus in immunohistochemical (IHC) studies of IGF-1R in human tumors, confounding attempts to assess the predictive and prognostic significance of IGF-1R expression and subcellular localization. Likely sources of variation include use of different IGF-1R polyclonal antibodies and methods for IHC. Here, we aimed to develop a robust IGF-1R IHC protocol using a monoclonal antibody, suitable for use in formalin-fixed paraffin-embedded (FFPE) tissues.

Methods: Using controls including samples of FFPE tissues and tumor cells of defined IGF-1R expression, we used IHC and western blotting to compare polyclonal antibody #3027 with monoclonals #9750 and #14534 (Cell Signaling Technology).

Results: Compared with #3027, the monoclonals exhibited superior discrimination between IGF-1R-high and IGF-1R-deficient cells in manual IHC, signal generated by #9750 reflecting differences in IGF-1R expression detected by western blotting. In tissues, IGF-1R detected by #14534 was predominantly plasma membrane-associated, while #9750 detected IGF-1R in the plasma membranes, cytoplasm and nucleus of prostate and renal cancers, recapitulating appearances we described using previous lots of #3027, and reflecting subcellular localizations reported using other techniques. Use of #9750 and #14534 in an autostainer showed adequate differentiation of high vs low IGF-1R cells, but did not recapitulate appearances of manually-stained tissues. We provide a detailed protocol for the preferred manual method using #9750.

Conclusion: Standardization of IGF-1R IHC will promote understanding of the role of IGF-1R in tumor biology, and its potential as a candidate prognostic and predictive biomarker.

Keywords: Cancer; Insulin-like growth factor; Type 1 IGF receptor; IGF-1R; Nuclear IGF-1R; Immunohistochemistry; FFPE; Antigen retrieval

Abbreviations: IGF: Insulin-like growth factor; IGF-1R: Type 1 IGF receptor; IHC: Immunohistochemistry; FFPE: Formalin-fixed, paraffin-embedded; NSCLC: Non-small cell lung cancer

Introduction

The contribution of insulin-like growth factors (IGFs) to cancer biology has been extensively studied in cell lines, revealing that IGFs activate type 1 IGF receptors (IGF-1Rs) to promote cell cycle progression, cell survival, motility and invasion [1,2]. These findings provoked interest in studying IGF-1R expression in clinical cancers, and in development of drugs that block IGF signaling. However, there has been striking variation in reported IGF-1R expression in tumors and normal tissues when detected by immunohistochemistry (IHC). For example IGF-1R was reported to be unchanged or downregulated in prostate cancer compared with benign prostate [3,4], although since our report of IGF-1R up-regulation at the mRNA and protein level [5] most publications support up-regulation [6-8].

This lack of consensus also confounds attempts to interpret results of clinical trials of novel IGF inhibitory drugs. Early trials reported striking clinical responses to IGF-1R inhibition e.g., Ref. [9], but later trials showed very limited activity in unselected patients (reviewed in Ref. [10]. This raises the question as to whether tumor IGF-1R expression correlates with sensitivity to IGF-1R inhibition. Preclinical reports supporting such a link include studies in non-small cell lung cancer (NSCLC), breast and colorectal cancer, rhabdomyosarcoma, Ewing family tumours and neuroblastoma [11-14]. However, other preclinical and some clinical studies found IGF-1R expression not to associate with response to IGF-1R inhibition in breast cancer, NSCLC and sarcomas [15-17]. While it is possible that IGF-1R is predictive of response in some tumor types but not others, it is also likely that technical differences in IHC protocols contribute to the apparent variation in significance of IGF-1R expression. Notably, the study conducted by Schwartz and colleagues found no difference in clinical activity of IGF-1R antibody cixutumumab with temsirolimus in patients whose tumors were IGF-1R ‘positive’ or ‘negative’ by IHC using the automated Ventana platform [17]. This trial also assessed tumor IGF-1R in a subset of patients by western blotting of fresh tumor lysates, resulting in detection of IGF-1R in all ‘IGF-1R negative’ tumors that were tested.

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Further complicating interpretation of IGF-1R IHC is the phenomenon of IGF-1R nuclear translocation, reported by several groups including ours [18-21]. In FFPE samples of human tumors, nuclear IGF-1R has been shown to associate with adverse prognosis in renal cancer [19], and with response to IGF-1R antibody in patients with sarcoma [21]. Lack of detection of nuclear IGF-1R prior to adoption of heat-based antigen retrieval [5,19] suggests that this also is influenced by technical variation in IHC methods.

These findings highlight the insensitivity of some IHC protocols, and the difficulties inherent in assessing the biological and clinical significance of IGF-1R expression and subcellular localization. Likely sources of variation include use of different antigen retrieval protocols and IGF-1R antibodies, and lot-to-lot variation in polyclonal antibodies. These technical factors may confound attempts to understand the role of IGF-1R in tumor biology, and the assessment of IGF-1R as a candidate prognostic and predictive biomarker. The aim here was to develop a robust IGF-1R IHC staining protocol that can be used to address these questions.

**Materials and Methods**

We used formalin-fixed paraffin-embedded (FFPE) human tumors that were surplus to diagnostic need, available in the Department of Cellular Pathology, Oxford University Hospitals NHS Foundation Trust, John Radcliffe Hospital, Oxford, and accessed under National Research Ethics approved study 07/H0606/120. As cell line controls, we used human MCF7 breast cancer, SKUT-1 leiomyosarcoma, PC3 prostate cancer, and A375M melanoma. IGF-1R depletion was induced by transfection with IGF-1R siRNA (Hs_IGF1R_1, Qiagen), with Allstars siRNA (Qiagen) as non-silencing control, as described [22]. Subconfluent cultures were scraped into ice-cold phosphate buffered saline (PBS), pelleted by centrifugation (1250 rpm for 5 minutes, 4°C) and resuspended in 10% neutral buffered formalin (NBF) for 1 hour at 23°C or overnight at 4°C. After centrifugation (1250 rpm, 5 min) the cell pellet was resuspended in 2% w/v formalin-agaroze (Sigma A9414 agarose, melting temperature 87°C, gelling temperature 36°C, in NBF) equilibrated to 60-65°C. After chilling on ice for 5-10 minute the cell pellet was embedded in paraffin, from which 1 mm cores were removed and inserted into a recipient block using a tissue microarrayer (Beecher Instruments, Sun Prairie, WI, US), and annealed at 45-50°C for 20 min. Parallel cultures were lyzed for western blotting using IGF-1R antibodies and β-tubulin antibody (Sigma-Aldrich), as described [19]. IGF-1R IHC was performed on freshly-cut 4 µm tissue and tissue microarray (TMA) sections using #3027, #9750 and #14534 antibodies (Cell Signaling Technology). Table 1 shows the manual IHC protocol, which used methods based on Ref. [19,23], and Table 2 the automated protocol.

| Step                        | Protocol                                                                 |
|-----------------------------|--------------------------------------------------------------------------|
| Dewax, rehydrate            | Incubate for 8 min in Citroclear (TCS Biosciences Ltd, UK), 8 min Citroclear, 2 min 100% ethanol, 2 min 95% ethanol, 2 min 80% ethanol, 2 min 70% ethanol, 2 min 50% ethanol, 10 min distilled water (dH2O) |
| Antigen retrieval           | Buffer: 50mM Tris-Cl, 2 mM EDTA pH 9                                    |
| Wash                        | Wash slides in PBS 3 × 5 min                                             |
| Block endogenous peroxidase | Incubate slides in 3% H2O2 in dH2O at 23 °C for 10 min                  |
| Wash                        | Wash slides in PBS 3 × 5 min                                             |
| Block                       | Incubate slides in 5% BSA/5% goat serum in PBS 1 hr at 23°C. Drain blocking solution with Kimwipe (do not wash away) |
| Primary antibody            | Mark around tissue area with PAP pen. Place slides in humidified chamber. Incubate with IGF-1R-β antibody #9750 (Cell Signaling Technology), diluted 1:50 in blocking solution (5% BSA/5% goat serum in PBS), using 200-500 µl antibody solution per slide depending on tissue area. Incubate on level surface at +4°C overnight. |
| Wash                        | Wash slides in PBS 3 × 5 min                                             |
| Detection of bound antibody | Use MP-531-M3R25, Menarini Diagnostics (Winnersh-Wokingham, Berkshire UK). Add 1-3 drops (depending on area to cover) of rabbit probe (neat as supplied) for 15 min (=5 min to come to –23°C plus recommended 10 min). Wash in PBS 3 × 5 min, incubate with rabbit HRP polymer (neat as supplied) for 15 min at 23°C. |
| Wash                        | Wash slides in PBS 3 × 5 min                                             |
| Color development           | Incubate with DAB substrate (Envision) for 5-7 min                      |
| Wash                        | Wash slides in tap water 3 × 2 min                                       |
| Counterstain                | Counterstain with Mayer’s haematoxylin (Vector Laboratories, USA) for 40 sec |
| Wash                        | Wash slides in copious tap water until water runs clear                  |
| Dehydrate                   | Incubate slides for 2 min in 50% ethanol, 2 min 70% ethanol, 2 min 80% ethanol, 2 min 95% ethanol, 2 min 100% ethanol, 5 min in Citroclear |
| Mount                       | Mount using DePex reagent (VWR International, UK)                        |

Table 1: Protocol for preferred manual IGF-1R IHC. This is a modification of the protocol reported in Ref. [19,23] using IGF-1R antibody #9750, and detecting bound antibody with reagents from MP-531-M3R25, Menarini Diagnostics (Winnersh-Wokingham, Berkshire UK).

**Results and Discussion**

In our initial study of IGF-1R expression in prostate cancer, we had used polyclonal antibody sc713 (Santa Cruz) [5]. In a later study [24], sc713 was insufficiently specific, and we re-optimized our IHC protocol to another polyclonal, #3027 (Cell Signaling Technology) [19,24]. This antibody performed acceptably in western blotting (e.g., Ref. [19,25]), but we recently questioned the specificity of current lots of this antibody in IHC. Therefore, we tested currently available lots of #3027 in IHC using FFPE sections of MCF7 and SKUT-1 cells, which express high and low IGF-1R respectively. Using the manual IHC protocol, there was detectable IGF-1R staining in both MCF7 and SKUT-1 cells, signal in the latter suggesting lack of specificity (Figure 1A). In comparison, monoclonal antibody #9750 showed greater differentiation between SKUT-1 and MCF7 cells. We also compared #3027 and #9750 by western blotting of cancer cell lysates (Figure 1B). Probing of duplicate membranes confirmed that both antibodies detected immunoabsorptive IGF-1R of the predicted size: mature IGF-1R beta subunit of 98 kDa and IGF-1R pro-receptor of 220 kDa. We noted that #3027 generated stronger signal than #9750, paralleling differences seen in IHC (Figure 1A). IGF-1R expression was relatively high in MCF7, lower but clearly...
Protocol
Aquatex aqueous mountant (Cat no 3631235, VWR). Peroxidase block at A. Standard Bond wash with distilled water. Bondmax standard wash protocol with Bond wash solution (Cat no. AR9590). Leica Bondmax autostainer standard bake at 60°C for 30 min, followed by standard dewax and rehydrate program using Leica Dewax solution (Cat no. AR9222), 100% ethanol and deionized water. Block endogenous peroxidase Peroxidase block at 23°C for 5 min with Bond Polymer Refine kit (Cat no. DS9800). Wash Bondmax standard wash protocol with Bond wash solution. Primary antibody Diluted 1:50 in Bond antibody diluent (Cat no. AR9352) for 1 hr at 23°C. Wash Bondmax standard wash protocol with Bond wash solution. Detection of bound antibody Post Primary Antibody (5 min at 23°C) followed by standard wash and polymer from Bond Polymer Refine kit (Cat no. DS9800) for 8 min at 23°C. Color development DAB 10 min at 23°C from Bond Polymer Refine kit (Cat no. DS9800). Wash Standard Bond wash with distilled water. Counterstain Haematoxylin 5 min from Bond Polymer Refine kit (Cat no. DS9800). Wash Standard Bond wash with distilled water. Mount Aquatex aqueous mountant (Cat no 3631235, VWR).

Table 2: Automated IGF-1R IHC protocol. This protocol was tested on a Leica Bondmax autostainer to evaluate antibodies #9750 and #14534. The method produced acceptable specificity and sensitivity in cell controls (Figure 1D) but generated very weak staining in FFPE tissue (see Figure 2A) and is not recommended for future studies.

| Step | Protocol |
|------|----------|
| Bake, Dewax, rehydrate | Leica Bondmax autostainer standard bake at 60°C for 30 min, followed by standard dewax and rehydrate program using Leica Dewax solution (Cat no. AR9222), 100% ethanol and deionized water. |
| Antigen retrieval | Leica ER2 buffer pH 9 20 min 100°C |
| Wash | Bondmax standard wash protocol with Bond wash solution (Cat no. AR9590) |
| Block endogenous peroxidase | Peroxidase block at 23°C for 5 min with Bond Polymer Refine kit (Cat no. DS9800) |
| Wash | Bondmax standard wash protocol with Bond wash solution |
| Primary antibody | Diluted 1:50 in Bond antibody diluent (Cat no. AR9352) for 1 hr at 23°C |
| Wash | Bondmax standard wash protocol with Bond wash solution |
| Detection of bound antibody | Post Primary Antibody (5 min at 23°C) followed by standard wash and polymer from Bond Polymer Refine kit (Cat no. DS9800) for 8 min at 23°C |
| Color development | DAB 10 min at 23°C from Bond Polymer Refine kit (Cat no. DS9800) |
| Wash | Standard Bond wash with distilled water |
| Counterstain | Haematoxylin 5 min from Bond Polymer Refine kit (Cat no. DS9800) |
| Wash | Standard Bond wash with distilled water |
| Mount | Aquatex aqueous mountant (Cat no 3631235, VWR) |

Image 1: IGF-1R detection using polyclonal and monoclonal IGF-1R antibodies in human cancer cells. A. Manual IHC was performed on IGF-1R expressing MCF7 and IGF-1R deficient SKUT-1 cells using polyclonal antibody #3027 at 1:200 or monoclonal #9750 at 1:50 dilution. Scale bar 20 µm. B. A375M cells were untransfected (Nil) or transfected with control or IGF-1R siRNA. Other cell lines were untransfected. Whole cell lysates (50 µg) were analysed by western blot using 1:1000 dilution of #3027 (upper) or #9750 (lower), showing detection of IGF-1R β (98 kDa) and IGF-1R pro-receptor (220 kDa). Membranes were re-probed for β-tubulin as loading control. Dotted line: one lane removed. C. Manual IHC on control TMA using #9750 antibody at 1:50, showing representative appearances of FFPE cell pellets made from parallel cultures to those used in B. Scale bar 20 µm. D. IGF-1R IHC on FFPE sections of MCF7 and SKUT-1 cells using #9750 or #14534 at 1:50 in manual (upper) or automated (lower) IHC. Scale bar 20 µm.

Having confirmed the specificity of both monoclonal antibodies in control cells of known IGF-1R expression, we performed testing in FFPE samples of human cancer. Staining of prostate cancer using manual and automated IHC showed negligible signal using either antibody at 1:200 dilution, and detectable signal at 1:50 that was more intense in the epithelial than stromal components, as previously noted [5]. The two antibodies showed different staining patterns, with faint membrane and more marked intracellular staining by #9750, and pronounced membrane staining by #14534. In both cases, the staining was much fainter in the auto-stained slides (Figure 2A). Finally, we analysed IGF-1R expression and subcellular localization in clear cell renal cancer, in which we had detected prominent nuclear IGF-1R in our previous study [19]. At low power it was apparent that #9750 gave stronger nuclei staining than #14534 (Figure 2B, left). As in prostate cancer (Figure 2A), #14534 showed prominent membrane IGF-1R, while #9750 showed membrane and cytoplasmic signal. In #9750-stained tumor there was clear detection of nuclear IGF-1R that showed cell to cell variation, recapitulating appearances generated by previous lots of #3027 in our prior study (Figure 2B, upper center and right panels; compare Figure 4B panel H in Ref. [19]. In contrast nuclear IGF-1R was only faintly detected by #14534 (Figure 2B lower). Givn that nuclear IGF-1R is detectable by a range of methods including subcellular fractionation, immunofluorescence, proximity ligation assay and chromatin immunoprecipitation, and is reported to have prognostic and predictive significance [18-21], we suggest that #9750 is preferable to #14534 when using IHC to study IGF-1R expression and subcellular localization in human tumors.

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In conclusion, we developed a specific and sensitive manual IHC protocol using antibody #9750. This is a monoclonal antibody, ensuring consistent antibody supply in future work. Nuclear IGF-1R was only weakly detected by antibody #14534, and auto-staining did not recapitulate appearances of the manual #9750 protocol, likely reflecting differences in antigen retrieval and detection. We provide a detailed protocol for the preferred manual IGF-1R IHC method that we commend for future studies. Adoption of a uniform protocol should reduce inter-study variation and may help to clarify whether responses to anti-IGF drugs associate with IGF-1R expression or subcellular localization.

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